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FORENSIC MEDICINE  
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# FORENSIC MEDICINE

AND

## TOXICOLOGY.

BY

J DIXON MANN, M.D., F.R.C.P.

**SIXTH EDITION, REVISED THROUGHOUT**

BY

**WILLIAM A. BREND, M.A., M.D., B.Sc.,**  
OF THE INNER TEMPLE, BARRISTER AT LAW, LECTURER ON FORENSIC MEDICINE  
AND TOXICOLOGY, CHARING CROSS HOSPITAL

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## PREFACE TO THE SIXTH EDITION.

THE fifth edition of this work was quickly exhausted notwithstanding the years of war, and for about eighteen months copies have not been procurable. The present edition has been revised throughout and brought up-to-date, whilst to avoid undue extension the work has been reset upon a larger page. The chapter on the forms of insanity has been to a large extent re-written. Additions have been made relating to cremation, and to poisoning by salvarsan, tetrachlorethane, trinitrotoluene, and gases of warfare. The section on food poisoning has been modified in accordance with recent views as to the part played by acute bacterial infections in conditions formerly ascribed to ptomaine poisoning.

I am indebted to Professor Harvey Littlejohn of Edinburgh for kind permission to reproduce his excellent photographs of the effects of decomposition in water.

W. A. B

THE MEDICAL SCHOOL,  
CHARING CROSS HOSPITAL, W C  
*March, 1922*

## PREFACE TO THE FIRST EDITION.

THIS work has been written chiefly as a Text-Book for Students of Medicine, it is hoped that it may also prove useful to Practitioners and others who are interested in the subject of Forensic Medicine. Since Medical Practitioners in general acquire much of their expert medico-legal knowledge from the study of reported cases, no pains have been spared in the selection, from a wide field of English and Foreign periodical literature, of typical examples illustrative of the subjects dealt with.

The section on Toxicology has been arranged with a view to simplicity and convenience of reference rather than to the attainment of an ideal classification.

I have to thank my friend and colleague Professor A. H. Young for valued assistance and advice on anatomical and morphological subjects.

J DIXON MANN

OWENS COLLEGE, MANCHESTER,  
*April*, 1893



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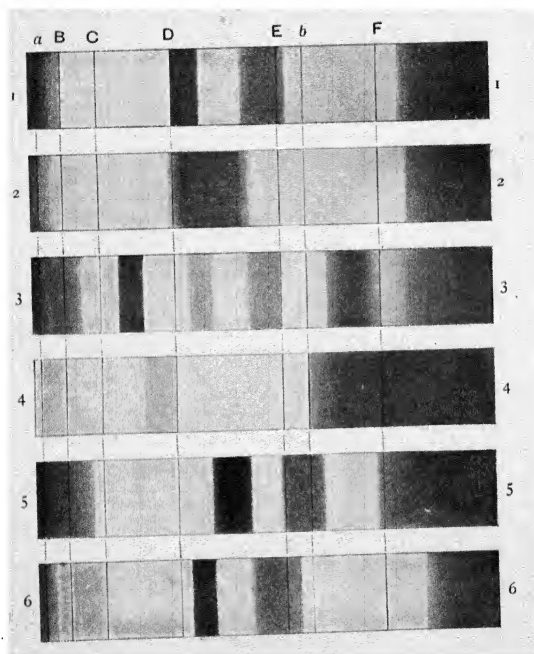
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# FORENSIC MEDICINE

AND

## TOXICOLOGY.

### PART I—FORENSIC MEDICINE.

#### CHAPTER I

#### INTRODUCTION—CORONER'S COURT—ASSIZES.

FORENSIC MEDICINE or Medical Jurisprudence is a many-sided subject, a knowledge of it demands more or less acquaintance with medicine in all its branches, and with the collateral sciences. It has been said that any medical practitioner who has a fair knowledge of his profession only requires common sense to qualify him as a medical jurist. This is a specious but fallacious statement. A knowledge of surgery, or of obstetrics, for example, limited to that required for the relief of suffering, would be of little use in unravelling many of the complex questions with which the medical jurist has to deal, although the branches of medicine named may be those which are appealed to for that purpose. Much of the knowledge and skill possessed by the surgeon is useless to the medical jurist, and that knowledge of a surgical character which is important to the medical jurist is all but useless to the surgeon. Take the case of a wounded man—the surgeon examines the wound in order to determine the best treatment for its cure, to him it is of minor moment as to whether the wound was self-inflicted or not, or as to whether it might have been produced by a certain weapon, or could not have been thus produced. The attention of the medical jurist, on the other hand, is chiefly directed to these points. Something more is required for forensic purposes than a knowledge of the various departments of medicine in relation to the healing of the sick, and this additional knowledge cannot be replaced by common sense, it necessitates special training. If a medical man were to commence his career as a medical jurist with no other preparation than a knowledge of medicine and surgery and the allied sciences solely as regards the art of healing, without being acquainted with the connecting links by which they are made subservient to forensic practice, he would inevitably overlook those features of a case which, correctly interpreted, yield the required information.

But while forensic medicine is a subject which requires special study, it is unfortunately the case that the opportunities for such study in this country

are limited. The applications of medical jurisprudence occur most frequently in connection with coroners' inquests. The inquiries in court are open to the public, but there is no right for any unauthorised person to be present at the post-mortem examinations. Nevertheless a student who wishes to gain a practical knowledge of forensic medicine should seize opportunities of attending inquests, and he will often find it possible to obtain permission to be present at autopsies. In France and Germany the judicial authorities afford special facilities for the training of medical students in this subject.

A word must be said as to the scope of forensic medicine. It has been defined as "the application of medical knowledge to the purposes of the law." This definition relates chiefly to such matters as the investigation of injuries, post-mortem examinations, and toxicological analyses in cases where crime has possibly been committed, the examination of either the living or the dead where questions of compensation are involved, and examinations in suits of nullity of marriage or cases of insanity. But it is highly desirable that every medical man should also have some knowledge of the statutes which directly affect him in the exercise of his profession, and these are now frequently included in the teaching of forensic medicine. For example, a practitioner should have at least a general knowledge of the constitution and powers of the General Medical Council, and should be aware both of the privileges and of the obligations he acquires by the act of registration. He should know his duties in regard to the certification of deaths, the notification of births, and the notification of infectious and industrial diseases. A timely knowledge of his responsibility in the treatment of his patients and of the obligation to maintain secrecy in professional matters may save him an unpleasant experience in the law courts. All these subjects are fully considered in Chap. XXV.

In some directions it is not easy to draw a dividing line between forensic medicine and the science of hygiene or public health. Such matters as the adulteration of food-stuffs, the prevention of venereal disease by the State, and restrictions on the sale of patent medicines, though primarily directed towards improvement of the public health, involve questions of law, and of the liberty of the individual, which justify their being brought under the purview of the medical jurist. Together these two branches constitute what is known as State Medicine, which may be regarded as embracing the entire range of activities connecting the State with the science and practice of medicine.

As knowledge of forensic medicine is required most frequently in the courts of law, it is convenient to begin with a brief description of the procedure observed therein.

### THE CORONER'S COURT.

The office of King's Coroner or "Crownor" is one of the oldest in the country, the first authenticated reference to it being contained in a precept to the justices in Eyre in the year 1194. The coroner was in early times a revenue officer of the Crown, and was charged with the duty of finding out criminals, extorting confessions if necessary, and confiscating their goods for the King. Until the time of Magna Charta he also held Pleas of the Crown, that is, acted as a criminal judge. Other duties were to take possession of treasure trove, wrecks, stranded whales, and royal sturgeons, and to assess the value of the *deadand*, that is the animal or inanimate object which had led to a death and was accordingly forfeited to the Crown. Thus if an ox gored a man to death, the ox or its monetary value was forfeited. As late as 1838 a coroner's jury levied a *deadand* of

£1,500 upon the boiler of the steamship *Victoria*, which had exploded and caused loss of life. One object of inquiring into the cause of a death was to ascertain whether or not there was property which escheated to the Crown, this being the case if the deceased was an outlaw or a felon, including, of course, a suicide.

But with the exception of that relating to treasure trove, these duties have become obsolete, and the Coroners Act of 1887 now merely requires that the jury shall certify how, when, and where the deceased came by his death. Very diverse interpretations are however placed upon these words by different coroners, and there is consequently a great range of variation in the principles and practice adopted in different courts. Some consider that their duty begins and ends in determining whether or not crime has been committed. In a case free from suspicion they may authorise a medical man to give a certificate even if he be unaware of the pathological cause of death, or where an inquest is held may be satisfied with a verdict of death from "natural causes." Others, for scientific or statistical purposes, inquire into deaths in which the pathological cause is unknown even if there is no reason to suspect violence. Some coroners consider that they are bound to hold inquests in all deaths under anæsthetics, others only if there are allegations of negligence. One coroner will scarcely hold an inquest without calling medical evidence, another only requires such testimony occasionally. The proportion of post-mortem examinations to inquests held ranges from over 90 to less than 20 per cent. The editor has elsewhere urged that in view of the importance of obtaining accurate vital statistics, the pathological cause of death should be fully ascertained in every case.

The circumstances under which a medical practitioner should notify the coroner of a death are considered in Chap. XXV. We are for the moment merely concerned with procedure. This is very simple. The jury having been sworn are required to "view" the body, which in most courts consists of their filing past a window on the other side of which is the deceased closely shrouded so as to conceal all but the face. The value of this proceeding is open to question. It is urged that it helps the jury to understand the medical evidence, but it is difficult to see how this can be the case without the condition of the internal organs being also shown. On the other hand, it undoubtedly leads to misconceptions. For example, the jury are very apt to regard post-mortem staining as the effect of violence, or to jump to the conclusion that emaciation indicates wilful neglect. Witnesses are examined on oath, their evidence is recorded, and in the event of further proceedings being taken, they are bound under a pecuniary penalty to appear at the superior court to which the case is transferred. The medical evidence is usually taken last, but in some courts the coroner provides the medical man with a form to fill up and return to him after making the post-mortem examination. He thus has the advantage from the beginning of the inquiry of knowing what the medical evidence will be. This is convenient, and if the form is properly drawn up has the further result of ensuring that the practitioner makes no omissions in his examination. There is no statutory obligation upon him to fill up the form, but it is usually desirable that he should do so. If the evidence is incomplete, and there is a prospect of additional evidence forthcoming, the coroner may adjourn the inquest. When a coroner's inquisition charges a person with murder or manslaughter, the coroner issues a warrant for the arrest of the accused person, unless he is already in custody. When the offence is manslaughter, the coroner may accept bail. As the proceedings are not directed against anyone—that is, no one is

on trial—it is not necessary that a suspected person should be present, though he is allowed to be so if he wishes

Under the Juries Act 1918, coroners are empowered to hold inquests without a jury. At present this is a temporary provision, but it may be retained permanently.

The medical witness should remember that almost all the criminal cases which attain the notoriety of *causes célèbres* are initiated in the Coroner's Court, and that the evidence he there gives may subsequently be subjected to the keenest scrutiny by subtle intellects specially trained for the purpose. The fact that the majority of the cases which come before the Coroner's Court end there, too often conduces to carelessness on the part of the medical witness in the preparation of the evidence he is about to give. Trusting to the impulse of the moment, he expresses opinions which he subsequently regrets having uttered. All statements there made are committed to paper in the form of depositions to which the witness appends his signature. Copies of these depositions are in the hands of both judge and counsel when the case comes before the assizes.

The law of evidence is not necessarily observed in the Coroner's Court, and the coroner may accept any evidence—hearsay or otherwise—that he likes. Counsel and solicitors are allowed to be present in the interests of clients, but have no *locus standi*. Theoretically they have no right to act as defenders or cross-examine witnesses, and can only put questions through and by permission of the coroner. In practice, however, considerable latitude is allowed in this respect, even statements which approach very closely to addresses to the jury being sometimes permitted. The Coal Mines Regulation Act of 1887 provides for the examination of witnesses by counsel representing persons whose interests are affected, subject nevertheless to the order of the coroner.

### MAGISTRATES' COURT.

Another preliminary court of inquiry is the **Magistrates' Court of Petty Sessions**. In this court the proceedings take the form of an investigation as to the culpability or non-culpability of a person accused of some act of criminality, or negligence of a criminal nature. As it is now a question of guilt or innocence, the accused person must be present, for the same reason witnesses may be examined and cross-examined by counsel, if no arrest has been made the magisterial investigation cannot take place. Unimportant cases—such as simple assaults—may be dealt with summarily. In cases of suspected manslaughter or murder it frequently happens that the magisterial investigation is held during or immediately after the coroner's inquest, and the medical witnesses who have there appeared are called upon to repeat their evidence before the magistrates. If the evidence is deemed sufficiently conclusive of culpability, the prisoner is committed for trial to a superior court, the witnesses being bound over to appear there and give evidence. The summons to attend at the assizes is called a *subpoena*, which, when tendered with reasonable travelling expenses, every witness in a criminal case is bound to obey. The relative obligations of common and expert witnesses will be subsequently discussed.

### ASSIZES.

The **Assizes** are courts in which both criminal and civil cases are tried. Usually two judges are present: one presides over the **Crown Court**, where the

The first of the permanent teeth appear on ground previously unoccupied—behind the rearmost of the temporary set. As a rule, they appear before any of the temporary teeth are lost, so that, on counting the teeth in one jaw from the central line backwards, if there are only five, they belong to the temporary set; if there is a sixth, it belongs to the permanent set. The rest of the permanent teeth replace the temporary teeth in the same order in which the temporary teeth appeared. The permanent molars appear at intervals of about six years. A child, nine years of age, will have twelve permanent teeth; at thirteen or fourteen, it will have twenty-eight—that is, all except the four wisdom teeth.

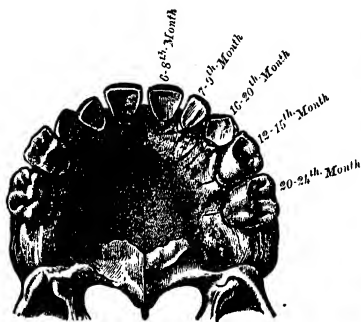


Fig. 4.—Temporary teeth (upper jaw).  
(Macalister's *Human Anatomy*.)

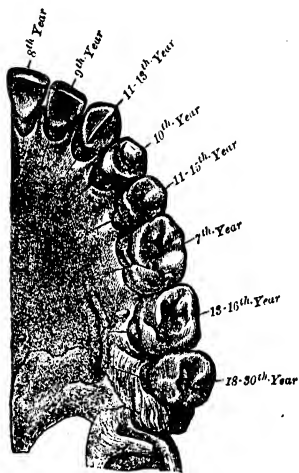


Fig. 5.—Permanent teeth.  
(Macalister's *Human Anatomy*.)

In advanced life, the bones of the head tend to become thinner from absorption of the diploe. The long bones become lighter and more fragile, the inorganic components being in excess. Many of the teeth are absent, and those which remain are worn down and discoloured.

The following Tables are arranged (mostly from Quain's *Osteology* by Thane) with the view of showing the probable age of a body or skeleton up to puberty, from the presence of the principal centres of ossification, and to later adult age from the union of epiphyses with bones, or of one bone with another:—

## CHAPTER II

**MEDICAL EVIDENCE : ORAL AND DOCUMENTARY.**

**Evidence** may be of two kinds (1) Evidence as to facts that have come under the observation of the witness (2) evidence as to the interpretation of facts, founded on a knowledge possessed by the witness of a special subject. Witnesses of the first kind are called "common witnesses", those of the second are known as "expert witnesses". The term "skilled witness" sometimes met with is best restricted to a witness who has special knowledge of the subject under investigation, such as a medical man or engineer, whether he be giving common or expert evidence.

**The Common Witness,** or witness to facts.—A medical man acts as a common witness when he gives evidence as to the condition of a wounded person examined by him. In such a case his duty is to describe the nature of the wounds, the general condition of the patient, and other circumstances that he observed at the time he made the examination. A common witness is obliged to give evidence if legally summoned to do so.

**The Expert Witness.**—When giving evidence solely as an expert, the witness acts as an interpreter of facts without having personal knowledge of them, usually a medical witness acts both as a common and as an expert witness, his skilled or expert opinion being founded on facts that he himself has observed. To continue the illustration above given of the duties of a common witness.—After describing the actual condition of the wounded person, he may be asked if the wound was of such a nature as to be dangerous to life. In answering this question the witness is no longer giving evidence to facts—he is acting as an expert, and is thus combining the functions of a common and of an expert witness. Before giving purely expert evidence—that is, evidence founded on facts of which he has no personal cognisance—it is necessary that the witness should have heard the facts on which he expresses an opinion stated on oath before the court.

The obligations of the expert witness are not so easily defined as those of the common witness. Hypothetically the knowledge possessed by an expert, who has no personal acquaintance with facts relating to a given case, is his own property, and therefore he ought not to be obliged to part with it against his will. This view has been taken by more than one judge. Lord Campbell ruled that a scientific witness was not bound to attend on being served with a subpoena. Justice Maule ruled that an expert is under no obligation to give evidence before a court of law. Unfortunately for experts, the difference of opinion with which they are proverbially accused pervades the judicial bench. Other judges have ruled that wilful neglect of a properly served subpoena constitutes contempt of court. In the face of decisions so adverse it is difficult to determine what the law on the subject really is. It is probable that an expert who had no personal acquaintance with a case might neglect a subpoena without rendering himself liable to attachment for contempt of court, but he might render himself liable to an action for damages. Such an action, though futile

so far as obtaining a verdict goes, would subject the defendant to much trouble and to some pecuniary loss in defending it. Having regard to the uncertainty of the results, the safest course would be to obey the subpoena under protest.

Previous personal knowledge of the facts of a case precludes a witness from taking any possible advantage of the *status* of an expert witness. There is no doubt as to the obligations of a witness so situated, he must obey a subpoena in his capacity as a common witness, although the evidence he is going to give may be of an expert character.

**Professional Secrets.**—It is an honourable law of the medical profession that confidential statements made by a patient to a medical adviser are held to be inviolable secrets. In a court of law this inviolability is overruled, a medical witness, if asked, is bound on the ruling of the judge to reveal any secrets that have come to his knowledge whilst in attendance on a patient. However repugnant it may be to the feelings of a medical man to violate the confidences of the consulting-room, he has no option. If, when in the witness-box, he refuses to obey the ruling of the judge and to answer a question involving the betrayal of a secret which is really the property of his patient—it having been revealed to him in trust and under the conviction of absolute confidence—he renders himself liable to committal for contempt of court. It is conceivable that a medical man might feel the obligation to secrecy so great as to compel him to decline to answer a question involving betrayal of the confidence of his patient. Such a step, however, should not be taken without a profound conviction of duty. A good citizen obeys the law, although he may have scruples in doing so, therefore, a witness should not set his private judgment against authority without very searching self-inquiry, an obstinate conviction must not be mistaken for a sense of duty. In the majority of cases it will probably be compatible with his sense of duty if the witness enters a protest against answering the question and then bows to the requirements of the law.

An interesting decision relating to the question of professional secrecy in the witness-box was given by Mr Justice McCardie in a case heard before him in January, 1920<sup>1</sup>. In this case the wife, who was petitioning for a divorce, alleged cruelty by the communication of venereal disease, and called a medical man who had attended her to prove that she had suffered from syphilis. The petitioner had received her treatment at a public Venereal Diseases Clinic. Article II (2) of the Public Health Venereal Diseases Regulations, 1916, lays down that "all information obtained in regard to any person treated under a scheme approved in pursuance of this article shall be regarded as confidential." In view of this regulation the medical man demurred to giving evidence. The judge, however, ruled that the evidence must be given.

When giving evidence, the witness may refresh his memory by referring to notes made by him at the time of, or immediately after, the events or proceedings to which he is testifying. Such notes must have been written by himself, and must be the original copy, he must not copy the original notes and use the copy in the witness-box, or, if he does so, he must keep the rough copy so that it may be compared with the transcript, the two must agree word for word. It is much better to use the original notes, and thus save explanation and discussion. In using notes in the witness-box the witness is only allowed to refer to them from time to time in order to refresh his memory for figures, dates, names of persons or of places, etc., he may not read his notes

<sup>1</sup> *Brit Med Journ*, Jan 17th and Jan 24th, 1920

in consecutive sentences. The law requires that the evidence tendered by a witness in court should be oral and not documentary.

In the course of a trial it is not unusual for a counsel to quote from a text-book on forensic medicine, or from some other book containing matter germane to the question at issue. If the witness disagrees with the quotation it is well for him to ask to be allowed to look at the book, possibly by reading the context he may find that the difference of opinion is only apparent, in any case, before accepting or denying a quotation, the meaning of which he does not fully comprehend, it is advisable for the witness to read the passage himself. He should also notice the date of publication. A book should not be quoted if the author is living. The objection is founded on the principle that evidence should be oral and delivered on oath in the witness-box, so that the giver of it may be cross-examined, this of course is impossible if the author is dead, and therefore his writings are admissible. The witness himself should not quote authorities to substantiate his opinions unless asked, the opinions he expresses are supposed to be the outcome of his own knowledge and experience.

Medical and expert witnesses are usually allowed to be present in court whilst the trial in which they are going to give evidence is proceeding. It is often absolutely necessary for the expert to become acquainted with the evidence tendered by the witnesses to fact, seeing that the opinions he forms are founded on this evidence. Merely having a transcript of the evidence delivered by the witnesses to fact read to him, in place of hearing it delivered is liable to lead to error of interpretation on the part of the expert. Further, if the evidence to a certain fact is lacking in some detail which is necessary to be brought out in order to arrive at a correct conclusion, the expert witness, if in court, has the opportunity of communicating with his counsel so that questions may be asked and the doubtful point cleared up.

A few words as to the giving of evidence before courts of law will be of assistance to the inexperienced medical witness. One of the most important points to remember is that, although the medical witness to a great extent deals with technical matters, he is addressing an audience which may be regarded as ignorant of technical terms. The judge and the counsel, although men of the highest intelligence and education, are not for the most part versed in the technical expressions by which medical men are accustomed to describe anatomical and pathological details. The medical witness has only to listen to lawyers discussing some point of law to appreciate the difficulty interposed to members of another profession by the use of technical words and phrases. If the judge and bar are not in a position to follow medical evidence couched in technical terms, much less are the jurymen—men of very ordinary intelligence and education—able to do so. The description of an injury which would be perfectly suitable if given before a medical society might be just as unsuitable if delivered in the witness-box. The avoidance of technical terms necessitates a good deal of paraphrasing, as such terms are necessarily the most concise in which the speaker can convey his thoughts. It is obviously more convenient to speak of the "peritoneum" than to say "the membrane that covers the bowels", but the latter expression would, and the former would not, convey information to the occupants of the jury-box. On the other hand, great care must be taken that the use of simple language does not lead to misunderstandings. In an actual case which turned upon whether a perforation of the intestines was due to a kick or appendicitis, the medical witness spoke of "the lining membrane of the intestine." He was understood by counsel to mean the mucous membrane, and confusion was caused until the error was detected.



Where there is the slightest possibility of ambiguity the best plan is to use the technical term first and then explain it, for example, "the peritoneum, that is the membrane lining the outer surface of the intestine"

Next in importance to the avoidance of technical expressions is the avoidance of superlatives and hyperbole. All evidence should be given with exactitude and without any attempt at dramatic effect, the style to aim at is a matter-of-fact style. Every object or condition should be described in the simplest terms consistent with perspicuity. As already stated, many of the questions asked require simply affirmative or negative responses. When a long string of such questions is asked there is often a tendency on the part of the witness to anticipate one or more of the succeeding questions which he assumes are coming, this draws upon him a sharp reproof from the counsel, who has planned his interrogations, and means to follow them out in his own way.

If a witness is doubtful about the answer to a question it is better to say frankly that he does not know than to seek to escape the difficulty by giving an ambiguous answer. When a counsel insists on 'yes or no' to a question that the witness feels cannot be properly so answered, he should ask to be allowed to qualify his answer, always avoiding such appeals unless the obligation of the oath he has taken to tell 'the whole truth' is in danger of being violated. No witness is required to answer a question if the answer would incriminate him.

All answers should be uttered in a sufficiently audible voice as to reach the judge and the jury, and sufficiently slowly as to enable the former to take notes. A witness should avoid a reserved or defiant manner, as though giving evidence under protest. Questions relating to facts should be promptly answered, those involving interpretation of facts demand caution. Too ready acquiescence should not be given to an apparently careless inquiry which half assumes the answer. A guarded reply to a question the answer of which involves a certain amount of discrimination is equally desirable as is a frank and open reply to a question relating to a fact observed by the witness.

## DOCUMENTARY EVIDENCE.

Documentary evidence includes **dying declarations** and **medico-legal reports**.

A **dying declaration** is a record of the evidence given by a person who is dying, and who believes that he is dying, from the results of an injury sustained, or a poison administered, at the hands of some person or persons. For a dying declaration to be valid it is necessary that the person who makes it should feel convinced that he is about to die. It is further necessary that this should be definitely stated in the written declaration. As the individual is incapable of writing down his deposition, it is taken by a second person, and this duty may devolve on a medical man. If the condition of the patient, though critical, is such as to permit of a short delay, the police should be informed, it is their duty to secure the attendance of a magistrate, who will take down the declaration. In such a case all that the medical practitioner has to do, if required, is to pronounce as to the mental fitness of the deponent to make the declaration. Should the condition of the dying person be such that there is not time to procure the attendance of a magistrate, the medical attendant himself must write down the declaration. In doing this the following rules should be observed — If possible, the attendance of one or more intelligent persons should be procured to act as witnesses, this is not necessary, but when important issues are at

stake it is advisable to omit no precautions that can possibly be taken. The medical man should then satisfy himself that the person making the declaration is convinced that he is about to die, and the declaration should commence with a statement to that effect. Such a statement must be unqualified. On one occasion the addition of two words was sufficient to invalidate a dying declaration, thus -- "No hope of my recovery at present", the last two words destroyed the expression of certainty that death was imminent, and the declaration was rejected at the trial. *The exact words uttered by the dying person should be written down.* No questions should be asked, except such as are necessary to clear up any obscurity. The declaration should be limited to a statement of what was done to the deponent at the time the injuries were inflicted, his own actions are not relevant. When completed, the declaration should be read over to the deponent, and, if possible, he should then append his signature. In any case, the witnesses present, together with the writer of the declaration, should sign it.

In some cases death occurs so rapidly, that even the medical attendant may not be able to take down the dying man's words in writing before he dies. The duty of the medical man in such cases is to listen to anything voluntarily said by the patient, and to take an early opportunity of writing down the exact words, and signing the statement. If others were present when the words were uttered, the statement should be read over to them, and their signatures should also be appended.

Considerable responsibility devolves upon medical men who are in attendance on persons who have been criminally injured and are in danger of death. No time should be lost in communicating with the police, as, apart from the occurrence of unexpectedly rapid death, the patient may become delirious or comatose.

If a person who had made a dying declaration recovers, the document ceases to have any legal force. The law only admits evidence when tendered on oath. In the case of a dying person an exception is made. It is believed that an individual who is convinced that he is about to die, will feel himself equally under an obligation to tell the truth, as though he was in the witness-box, and had taken the oath. If, after making a dying declaration, the deponent recovers, he reverts to his normal condition as regards the law, and must tender his evidence before the court in the usual way. Dying declarations are not admissible in civil cases.

/ **Medico-legal reports** in criminal cases are more used in Scotland than in England. A medical report is divisible into two sections -- (1) The result of the examination of the case, or, in other words, the facts that have been observed. (2) The deductions drawn from a consideration of such facts.

It is absolutely necessary in a well-ordered medical report that this division should be rigidly adhered to. Every fact should be recorded before the inferential part of the report is begun. Short of absolute inaccuracy, nothing vitiates a medical report so much as the mixing of facts with deductions, first state the facts and then the conclusions drawn from them. The legal authorities for whom the report is made may know nothing of the case, and they require to become acquainted with all the facts before they are in a position to appreciate the deductions drawn from them.

The facts to be observed in the living are limited to those which fall within the scope of medical observation. A good report, while containing all that is necessary, will be free from extraneous matter. Nothing should be included in a report that does not come under the personal observation of the reporter,

indirect or hearsay evidence is as much out of place in a report as in the witness-box. To avoid error of reading, all dates should be written in full. Every detail that can possibly have a direct bearing on the case should be noted, it is impossible to foresee what questions may subsequently arise. When making a report concerning a dead body, everything, both as regards the body and its surroundings, should be scrupulously noted. When the autopsy is completed, and consequently all the available facts are obtained, the inferential part of the report is to be undertaken.

If the report refers to a living person a careful study of the facts is to be made in order to arrive at a conclusion as to the way in which an injury has been inflicted, and whether the statements of the wounded person are or are not borne out by the actual condition of the injured parts. If it refers to a dead body the questions are: How was death caused? Did it immediately follow the infliction of the injuries? If not, what interval intervened? Were the injuries of such a nature as to incapacitate the deceased from moving after they were inflicted?—with other observations to which attention will be directed when the mode of conducting a post-mortem examination for medico-legal purposes is described.

The advice previously given as to oral evidence is equally applicable to documentary evidence. The language in which a report is drawn up should be free from technical terms and exaggerated expressions. The report should not be too long—when giving oral evidence in court the judge and the counsel will take care that the witness does not err in this respect, when writing a report he has unlimited scope, and is sometimes apt to be verbose. It is to be remembered that the writer of a report has not done with it when he has despatched it to its destination. Copies are placed in the hands of counsel, who will insist on an explanation of every ambiguous phrase, the longer the report the more likely are ambiguous phrases to occur, for it is in the inferential part that verbosity asserts itself.

### CHAPTER III

## LEGAL PROCEDURE IN SCOTLAND.

THERE are certain differences with regard to legal proceedings in Scotland as compared with England.

Public prosecutors are appointed by the Crown, who conduct criminal prosecutions in both higher and lower courts. The Lord Advocate and the Deputy Advocates take charge of cases which come before the High Courts of Justiciary, the Procurator Fiscal appears before the lower courts.

The first step in a criminal prosecution is taken by the Procurator Fiscal, who, on information supplied by the police or by private persons, makes such inquiries as satisfy him with regard to the necessity or not for legal proceedings. Any person who is supposed to know anything about the case is interrogated privately before the Sheriff, or to use the legal term, is “precognosed.” The examination is not made on oath, unless the witness is suspected not to be telling the truth. The evidence obtained is written down and forms the precognitions. The counsel for the accused, as well as the counsel for the Crown, has the power of precognosing the witnesses.

The Sheriff or Justice, before the preliminary examination has been conducted, liberates the accused or commits him for trial in accordance with the nature of the evidence obtained. If the accused is committed, the precognitions are forwarded to the Crown Counsel in Edinburgh, who have the power either to stop the proceedings or to send the accused before the High Court, or one of the Circuit Courts of Justiciary, or before the Sheriff with or without jury. The Courts of Justiciary correspond to the Courts of Assize in England. Should the case be sent for trial, the persons who have been precognosed, or such of them as the Crown Counsel select, are summoned by writ as witnesses. Neglect of such citation, unless sufficient cause be shown, is punishable by a fine of £5, and also by imprisonment, from which the offender is only released on expressing his regret before the Court and tendering bail to appear to give evidence.

Common witnesses are not allowed to be in court except when giving evidence—this applies to medical witnesses also who appear as witnesses to fact. By mutual consent of the opposing counsel, expert witnesses are generally allowed to remain in court. When an expert is giving his evidence, the other experts are usually required to leave the court. An expert witness who has been in court during the delivery of evidence by common witnesses, cannot be examined as a witness to facts.

In addition to the verdicts—"Guilty" and "Not guilty"—returned by the jury in England, the Scotch law includes a third—"Not proven", this verdict, as well as "Not guilty," is final, the accused cannot again be tried for the same offence. In England the jury must be unanimous, in Scotland a bare majority can return a verdict.

The Procurator Fiscal performs the duties undertaken by the Coroner in England, but without a jury. If a dead body is found, or a case of suspicious death occurs, the Procurator Fiscal, on being informed, has the power of directing a medical man to make an examination of the body, and to forward him a report dealing with the case, all such reports being certified by the reporter "on soul and conscience." If the medical examiner is satisfied with an external examination, he may certify to the Procurator Fiscal without making an internal examination. If a complete examination is requisite, the Procurator Fiscal issues a warrant to the medical practitioner who has already seen the case, and usually associates with him another practitioner of experience. The warrant is countersigned by the Sheriff or Justice, and empowers the holders of it to take charge of the body, and to make such examination as the law requires. The warrant also enables the inspectors to exclude improper persons from the room where the examination is being made. If, notwithstanding this authorisation, the relatives of the deceased refuse to allow the examination to take place, the authorities granting it (on being informed) will take steps to remove opposition. To ensure completeness of examination, the Crown Office in Scotland issues a form of instructions to medical inspectors, which contains elaborate directions for making the necropsy.

A medical practitioner, whether previously acquainted with a given case or not, cannot refuse to be precognosed if duly cited to that effect. Refusal is met by a further warrant and, in case of contumacy, by imprisonment.

## CHAPTER IV

## EXAMINATION OF THE DEAD BODY.

**Post-mortem Examinations for Medico-legal Purposes.**—There are several important points to be observed when making a medico-legal necropsy, over and above the requirements of ordinary pathological investigations

**External Inspection.** The examination should be made in daylight, colour changes are often invisible by artificial light. If the body is seen on the spot where it was first discovered, attention should be paid to the following points.—The exact posture in which it lies, the expression and colour of the face, the position of the hands, whether clenched or not, if clenched, they should be examined for any substance possibly grasped by them. The presence of any odour about the mouth or nostrils should be noticed. The fingers should be examined for cuts or wounds. The condition of the dress, if disordered, indicating a struggle, or if it is soiled or stained with blood. Attention should be directed to the ground on which the body lies, and to that immediately around it, for signs of struggling and for objects that may have dropped, as fragments of clothing, etc. Any discovery should at once be recorded in writing. The presence or absence of body-heat, of hypostases, of cadaveric rigidity, and of putrefactive changes are to be observed. When an exhaustive investigation of the body *in situ* has been made, it may be removed to some place convenient for further examination.

The clothes are now taken off, and any cuts or injuries sustained by the clothing carefully compared with the underlying surface of the body. Marks resembling bruises should be sponged so as to make sure that they are not due to dirt or other external stain. Indications for identification are to be sought for in surface marks—*nævi*, moles, tattoo-marks, cicatrices, external abnormalities or loss of fingers or limbs, absence of natural, or presence of artificial, teeth, colour of the hair, height, weight, sex, age, state of nutrition, and indications of social position or of occupation. In women and female children the presence or absence of the hymen, any signs of recent violence to the genital organs, together with the presence of foreign substances in any of the natural apertures of the body, should be ascertained.

If there are wounds, examine them carefully as to their length and depth and the structures divided or injured, whether they could have been self-inflicted, and the kind of weapon that could have produced them. Examine the neck for marks of strangulation. If there is a gunshot wound, look for blackening or tattooing of the surrounding skin, and also for blackening of the hand.

The **internal** examination must include all the organs and cavities of the body. If this is not done, the counsel for the defence may assume the presence of disease in an important organ which has not been investigated, or it may be necessary to have a second examination made to clear up a doubtful point which ought to have been settled by the first examination. A median incision should be made from the chin to the symphysis pubis, avoiding the umbilicus and any wounds which may be present on the body. The incision through the

scalp should be made from ear to ear over the vertex. The skin is then reflected backwards and forwards and the skull sawn through, the use of hammer and chisel being avoided for fear of producing a fracture of the skull or of causing one already existing to spread. If there are any penetrating wounds produced by cutting instruments or by firearms, ascertain their direction, and, in case they are not self-inflicted, try to form an opinion as to the relative position of the deceased and his assailant. When bones, cartilages, or intervertebral substances are injured, it is well to remove the injured parts and preserve them as evidence. Look carefully for any acute or chronic morbid changes in the organs, especially in cases of suspected poisoning, or when there is no gross traumatic lesion which would account for death. The vagina and the uterus are to be examined for signs of recent delivery and for mechanical injuries, or for injuries produced by the introduction per vaginam of caustic or irritant substances. The vertebral canal should be opened and the condition of the cord ascertained, if there is any reason for suspecting that it has been injured or is in any way concerned with the death.

The period at which death took place may be a question of great importance, therefore observe the condition of the stomach as to food. For example, a woman ate a hearty supper at 10 P.M., the following day her body was found with the throat cut. A man who was known to have visited the deceased the night before she was found dead was arrested on suspicion, he admitted the visit, but proved that he left the house before 11 P.M. At the autopsy the stomach was found empty, without any signs of vomiting having occurred, showing that after the supper at least two or three hours had elapsed before death took place—thus, taken in conjunction with the time at which the accused left the house, afforded strong evidence in his favour. The condition of the bladder may also afford a clue, if, in a case like the above, the bladder is found full of urine the inference would be that the deceased lived some time after going to bed.

#### AVERAGE WEIGHTS OF THE PRINCIPAL ORGANS IN THE ADULT BODY

Brain, male,	46 to 53 ozs	Stomach,	4½ ozs
„ female,	40 to 47 ozs	Liver, male,	48 to 58 „
Lungs, male (right),	24 ozs	„ female,	40 to 50 „
„ „ (left),	21 „	Pancreas,	2½ to 3½ „
„ female (right),	17 „	Spleen,	5 to 7 „
„ „ (left),	15 „	Kidneys (each),	4½ „
Heart, male,	10½ „	Suprarenals (each),	1 to 2 drms
„ female,	9½ „	Uterus,	7 to 12 „

**Cases of Suspected Poisoning.**—Several large glass jars, preferably new, but in any case thoroughly cleansed, should be provided. If they are furnished with glass stoppers so much the better, if not, some bladder or gutta-percha tissue should be obtained, which may be secured by string over the mouths of the jars. It is convenient to have a large dish—a photographer's square porcelain dish is the best—for placing the stomach in when opening it.

Before opening the body, examine the mouth and lips for injuries caused by a corrosive, and ascertain if there is any peculiar odour given off from the mouth. After making the primary incision through the abdominal parietes, again try if any special odour can be distinguished, and if so obtain corroboratory evidence from those who are present, the same proceeding should be adopted when the stomach and intestines are opened. When the abdominal cavity is opened, look for signs of inflammation of the peritoneum or of any of the viscera,

especially of the peritoneal aspect of the stomach. Then place a ligature round the lower end of the œsophagus, and a double one at the commencement of the duodenum. Divide the œsophagus above its ligature, and the duodenum between the two, and remove the stomach. On a dish, as already described, open the stomach along the small curvature, taking care that none of the contents are lost. The contents may be poured into one of the jars, and the inner coat of the stomach examined forthwith, its colour when first opened being noted. Search should be made with the aid of a lens for crystals, fragments of leaves, berries, and other parts of plants, and for particles of pigments (such as indigo), which are mixed with certain poisons— as arsenic when sold in small quantities, and strychnine in the form of vermin-killer. Any suspicious substances found should be carefully collected and examined under the microscope. The intestines, large and small, separately ligatured, are to be removed and treated in the same way. In the case of corrosive and irritant poisons, the œsophagus should also be removed, opened, and its internal appearance noted, the effects of the poison being traced from the mouth down the digestive tract as far as any can be observed. The presence or absence of solid fæces in the lower bowel is to be recorded.

The colour of the blood, its condition as regards fluidity, and the colour of the solid organs generally, should be observed. Indications of fatty degeneration in liver, kidneys, and heart, of injection, especially of the kidneys, and of ecchymoses must be looked for. In addition to the stomach and intestines with their contents, the liver, kidneys, spleen, as much of the blood as can be collected, with the contents of the urinary and gall bladders, should be severally removed and placed separately in appropriate vessels for analysis. It is well to remove the brain with any fluid that is present within the cranium, especially in the case of volatile poisons, and to preserve it as above described. In cases of chronic arsenical and mercurial poisoning some of the spongy bones— the bodies of the vertebræ, or part of the skull— should also be sent. The viscera named should be sent entire, not small portions of them, this is necessary in order to provide material for two independent analyses should they be ordered. All vessels should be closed so as to be as nearly air-tight as possible, the mouths are then covered with paper securely tied, the knot of the string being well coated with sealing-wax impressed with the private seal of the medical man who makes the examination. Labels should be attached to the jars and bottles, on each of which a description of the respective contents, with the name of the individual from whom they were derived, and the date of the necropsy, should be clearly written. Two lists of the jars and contents should be made, one being forwarded along with the jars to the analyst, or to the authorities who take charge of them meanwhile, the other being retained by the sender. The jars should pass through as few hands as possible, when feasible, the person who makes the post-mortem should himself deliver them to the analyst. They should be kept in a cool place, but no preservative should be added to their contents.

It sometimes happens that the most careful post-mortem fails to reveal to the naked eye morbid changes sufficient to account for the death. Affections of the nervous system, such as epilepsy, or some of the zymotic diseases, as well as alkaloidal poisoning, furnish examples. Under these circumstances the practitioner should be careful not to magnify natural appearances into morbid ones. The clinical history in such cases may be of the utmost value in determining the cause of death.

In some cases it is advisable that two practitioners should conjointly make

the post-mortem examination. In case of doubtful or of obscure indications, the advice and countenance of a colleague is advantageous, and the division of labour—one practitioner making the section and the other recording the results—adds to the completeness of the investigation and to the facility with which it is made. Every step should be accurately recorded at the time, or in the event of the examination being made by one medical man only, immediately after its completion. If the notes are made by a colleague they should be read over on the spot by the operator, and then signed by both medical men. If a medical man is implicated, he must not be permitted to be present, he may depute another medical practitioner to represent him at the necropsy, but his representative must not take any active part in the proceedings. In all cases in which a legal inquiry is likely to take place, the medical practitioner in charge should refrain from making an examination until he receives an order from the coroner to do so. When an inquest is going to be held, the dead body is technically in the possession of the coroner until he has issued his order for burial, and consequently it may not be interfered with without his permission. In other cases the Anatomy Act of 1832 (2 and 3 Wm IV c 75, sec 7) provides that the executors, or other party having lawful possession of the body, may permit an anatomical examination to be made.

**Exhumation**—When suspicion of foul play arises after the body of the supposed victim has been interred, the coroner and also the authorities at the Home Office may order the body to be exhumed and a medical inspection made. The medical man deputed to examine the body should be present at the exhumation, and should previously see that adequate provision is made for making a full investigation. A relative or friend of the deceased should be present at the exhumation in order to identify the body. When the interment has been recent an ordinary post mortem examination can be made, but if the body has lain long underground decomposition will be more or less advanced and the usual post mortem appearances destroyed. In such cases injuries to the bones, especially those of the skull, and in women the uterus (which resists putrefaction longer than the other soft organs) may afford valuable evidence. Most frequently exhumations are undertaken in cases of suspected poisoning, in such cases the stomach and intestines are to be removed—if recent, they should be ligatured as described in the directions for the ordinary examination, and placed in clean glass vessels well secured. The liver, spleen, and kidneys should be all removed. When the presence of a metallic poison is suspected, as mercury or arsenic, some of the bones should also be taken, the bodies of the vertebrae for example. If the interment was remote, so that the coffin is decayed, it is advisable in cases of mineral poisoning to remove a little of the surrounding earth for chemical examination. However far putrefaction is advanced, neither preservative fluid nor disinfectant must be used when making the post mortem, nor added to the parts removed. The stage of the putrefactive changes in relation to the length of time the body has been interred should be noted.



## CHAPTER V

**AGE IN ITS MEDICO-LEGAL RELATIONS.**

THE question of age in the living may come under the notice of the medical jurist in relation to criminal responsibility, marriage, fecundity, viability, rape, and personal identity. In the dead, in relation to infanticide, criminal abortion, and personal identity.

A child under seven years of age is held by law to be incapable of committing a crime, and, consequently, is exempt from punishment. Above that age, but below fourteen years, a child is still deemed irresponsible, unless proof of such a degree of intelligence is forthcoming as to show that he understood the criminal character of the act committed by him. An "infant" under the age of fourteen years is presumed by law to be incapable of committing a rape, and, therefore, cannot be found guilty of the crime, nor of an attempt to commit it. At and after the age of fourteen a youth is held responsible for his actions, but he does not attain the full privileges of an adult until he reaches the age of twenty-one years. It is not until he attains his majority (twenty-one years) that he can make a valid will (1 Vict c 26). The day of birth is included in computing the age, and, therefore, a valid will may be made on the day before the twenty-first "birthday," as the law does not recognise a division of time less than one complete day. The obligation to serve on a jury does not affect a man until he has reached his majority. In courts of law evidence may be given irrespective of age, provided that a sufficient degree of intelligence is manifested by the child when interrogated by the judge as to his or her capacity to understand the necessity of speaking the truth. The marriageable age in this country is fourteen years for the male sex and twelve years for the female.

In determining age in the living, no reliable criteria are available after adult life is reached, in the young, the teeth yield evidence up to the thirteenth or fourteenth year. General indications, of course, exist, but their variability—from idiosyncrasy, mode of life, personal attention, etc—is so great, that to estimate the age of a living person between the two extremes of life is little more than guess-work.

In the **dead**, the case is different. There are developmental signs which, when found, limit the age of the individual in both directions sufficiently narrowly to enable a fairly accurate estimate to be made. No single sign, as a rule, is determinative, but when several are found to be in accord the expert is warranted in giving a decided opinion.

The most reliable information in the later foetal months, and in the earlier years of life, is afforded by the ossification of the bones. The length and weight of the body during intra-uterine life afford important data, as do also the degree of development of the nails on the fingers and the toes, the size of the external ear, the presence or absence of meconium in the intestines, of lanugo on the skin, and of the pupillary membrane in the eye, and, in the male, the position of the testicles. It is not necessary for our present purpose to go further back than the sixth month of intra-uterine life, at this period viability may be said to begin. The dead body of a foetus that can be proved not to have reached

the sixth month of utero-gestation may be regarded as having been still-born, which would, of course, negative the charge of infanticide.

To avoid error, the period of utero-gestation in the following tabulation is stated in complete months, for example, of six months' duration, not in the sixth month. The latter mode of expression is ambiguous. Gestation of six months' duration means that the full period named has been accomplished. Gestation in the sixth month means any time from the commencement to the end of the sixth month and, therefore, may mean five months and one day.

### DEVELOPMENT OF THE FÆTUS FROM THE SIXTH MONTH TO THE FULL TERM OF UTERO-GESTATION.

**Six Months**—Length, 9 to 13 inches. Weight, 1 to 2 pounds. The head of the fœtus is large in proportion to the body. The insertion of the funis is considerably below the middle of the body. The skin is red and wrinkled, the underlying fat, which subsequently imparts rotundity to the body and limbs, is only now commencing to form. The body is covered with downy hair or lanugo, and also with a thin layer of vernix caseosa—a white substance consisting of sebaceous matter derived from the skin, mixed with epithelium and lanugo. The bones of the head are widely separated at the sutures, the anterior and posterior fontanelles being open. The sylvian fissure is formed. The precentral, inferior frontal, and intra parietal sulci of the cerebral cortex appear.<sup>1</sup> The eyebrows and the eyelashes are beginning to form. The eyelids are adherent. The pupillary membrane, which is formed in the third month, is present. The external auricle measures 16 to 24 millimetres.<sup>2</sup> The finger nails are forming, but are quite soft, the toe nails are less developed. The scrotum is smooth and empty. The testicles are on the psoas muscles, below the kidneys. In the small intestine there is a little mucoid secretion, which may be coloured with bile pigment. Centres of ossification are present in the os calcis, the manubrium, and in the bodies and laminae of the sacral vertebrae.

**Seven Months**—Length, 12 to 15 inches. Weight, 2 to 4 pounds. The skin is rather paler, and is well covered with lanugo and vernix caseosa. The lanugo is beginning to disappear from the face, that of the scalp is taking on the character of hair and is becoming darker. The superior precentral and the superior frontal sulci appear. The eyelids are not adherent. The pupillary membrane, which reaches its highest development during this month, begins to disappear. The external auricle measures 26 millimetres. The finger nails do not quite reach the ends of the fingers. The testicles are near the abdominal ring. Meconium is found in the large intestine. Examined microscopically, this substance is seen to consist of mucous corpuscles, epithelium from the intestine, small crystals of bilirubin (very like hæmatoidin crystals), crystals of stearic acid, and vernix caseosa. Centres of ossification are present in the first piece of the body of the sternum and in the astragalus.

**Eight Months**—Length, 15 to 17 inches. Weight, 4 to 5 pounds. The insertion of the funis is only slightly below the mid-point of the body. The skin is a little paler, and is more filled out by increased amount of fat beneath it. The face retains a wrinkled appearance. The lanugo is disappearing. The pupillary membrane has generally disappeared. The external auricle measures 26 to 28 millimetres. The nails feel harder, and have reached the ends of the fingers, but probably not the ends of the toes. The testicles are in the inguinal canal, or they may have reached the upper part of the scrotum, especially the left testicle. Valvulae conniventes are formed in the small intestine. The kidneys are now larger than the adrenals, and the bladder may contain urine. A centre of ossification is present in the second piece of the body of the sternum.

**Nine Months, at term.**—Length, 18 to 20 inches. Weight, 5 to 8 pounds. The head measures transversely  $3\frac{1}{2}$  to 4 inches, sagittally,  $4\frac{1}{4}$  to 5 inches. The shoulders measure  $1\frac{1}{2}$  inches across, the hips, 4 inches. The umbilicus is only three quarters of an inch below the mid point of the body. The skin has lost its rosy tint, and resembles more in colour that of the adult. The limbs and body are plump, and the face has lost its wrinkles. The lanugo has almost disappeared. Vernix caseosa is only present in quantity on the back and on the flexor aspect of the limbs. The hair on the head is mostly dark and is about an inch long. Along the lines of the sutures the bones of the skull are close together, but

<sup>1</sup> Cunningham, *Contribution to the Surface Anatomy of the Cerebral Hemispheres*, 1892.

<sup>2</sup> von Troeltsch, *Die Anatomie des Ohres*.

the parietal and occipital bones are only united by membrane and are freely movable. The posterior fontelle is closed, the anterior fontelle is not closed. The secondary sulci of the brain appear, and the surface is more highly convoluted. The eyelashes and the eyelids are well formed. The external auricle measures 3.3 to 3.6 millimetres. The cartilages of the nose and the ears feel hard. The nails project beyond the tips of the fingers, and they reach the tips of the toes. The testicles are in the scrotum, which is well corrugated. Meconium is present in the large intestine only. The breasts in both sexes are well formed, and contain some secretion. In the lower end of the femur there is a centre of ossification which measures 0.2 inch in diameter. This centre of ossification is of great importance to the medical jurist when investigating the development of the fœtus in cases of infanticide. It appears with tolerable constancy about a fortnight before full term, the epiphysis in which it is developed being the only one in which ossification begins before birth. Centres of ossification are also present in the cuboid, in the first coccygeal vertebra, and in the third piece of the body of the sternum.

The following table shows the more important developmental changes which take place in the fœtus from six months to full term --

TABLE OF DEVELOPMENTAL CHANGES IN THE FŒTUS

Months	Length in inches	Weight in pounds	Nails	Pupillary Membrane	Testicles	Centres of ossification
6	9-13	1-2	Forming	Present Eyelids adherent	On psoas muscles below kidneys	Os calcis Manubrium Bodies and laminae of sacral vertebrae
7	12-15	2-4	Not reached ends of fingers	Partly present Eyelids open	About abdominal rings	First piece of body of sternum Astragalus
8	15-17	4-5	Ends of fingers, but probably not of toes	Disappeared	Inguinal canal, may be in scrotum Left often before right	Second piece of body of sternum
9	18-20	5-8	Project beyond tips of fingers, reach extremities of toes		Usually both in scrotum, which is corrugated	Cuboid Third piece of body of sternum First coccygeal vertebra Lower epiphysis of femur

The above criteria of the stage of development of the fœtus are subject to considerable variation, and consequently a decided opinion should not be given unless, in an individual case, several of the most important reasonably coincide. As regards dimensions and weight, it is to be remembered that male children usually exceed female children in both respects. The variation in weight of children at term is considerable. Ortega<sup>1</sup> delivered a woman of a still-born child which measured 27 inches in length and weighed 24½ pounds, it measured 7½ inches across the shoulders. Playfair<sup>2</sup> quotes the case of a child born of parents of gigantic stature—the mother being 7 feet 9 inches in height and the father 7 feet 7 inches. The child was still-born, it measured 30 inches in length and weighed 23½ pounds. The weight of the fœtus at term, being

<sup>1</sup> *Nouv. Arch. d'Obstét. et de Gynéc., 1891*

<sup>2</sup> *Sc. and Prac. of Midwifery, 1893*

influenced by the state of nutrition, is less constant than the length. Rickets retards ossification, and thus interferes with the closure of the fontanelles and with the development of the various centres of ossification normally existing at term.

In the newly-born, putrefactive changes take place with great rapidity and seriously interfere with the indications afforded by the weight, the condition of the skin, hair, nails, cartilages of the nose and ears, and, by producing opacity of the cornea, with evidence afforded by the pupillary membrane. It is to be observed that the absence of the pupillary membrane does not in itself determine the maturity of the fœtus, nor is the absence of the testicles from the scrotum to be accepted as an indication of immaturity, they not unfrequently remain in or above the inguinal canal until or beyond puberty. The presence or absence of the centre of ossification in the lower epiphysis of the femur, although not infallible, is perhaps the most reliable means of enabling an opinion to be formed as to the maturity or otherwise of a fœtus even when putrefactive changes are so far advanced as to render other indications valueless. It is to be remembered that the difference between a fœtus at eight months and one at full term is not sufficiently pronounced as to enable a positive opinion to be given. This, however, does not affect the issue in the greater number of cases that come before the medical jurist, seeing that in either case the child might have been born alive.

**After birth,** the circulation soon becomes of the adult type. The cuticle begins to desquamate about the third day. The skin, at first hyperæmic and red in colour, subsequently acquires a yellowish tinge. When desquamation is finished— the time occupied varying from one to two weeks or more, according to the vigour of the child— the skin assumes its permanently normal colour. During the first three days the child loses weight. A few hours after birth the intestines relieve themselves of the meconium which accumulated during intra-uterine life. At birth, the normal umbilical cord is plump, spiral, and of an opal colour, in weakly children it is flaccid and is much thinner. After division of the cord, the portion attached to the child becomes flabby and begins to shrivel up on the first or second day, desiccation commences at the free end, advances towards the point of insertion, and is completed about the third or fourth day. The cord is then flattened, and has a parchment-like, translucent appearance, exhibiting the arteries and vein as red lines. If a cord that has undergone mummification is soaked in water, it does not return to its previous condition. About the fourth or fifth day the cord separates by ulceration close to the abdominal wall. Around the point of detachment is an inflammatory zone, which persists some time after separation. The usual purulent secretion which accompanies ulcerative processes is more or less present. The degree of, and the time occupied by, the inflammatory processes which attend separation of the cord vary with the development and vital activity of the child, the feebler the child, the longer the process of detachment. Cicatrization of the navel is usually completed in about eight to twelve days.

After separation of the funis and completion of desquamation of the cuticle, the progress of dentition and the development of the various centres of ossification that form after birth afford the most reliable criteria of age. There are, however, other indications which may be of use.

**The capacity of the stomach** rapidly increases after birth. Ashby and Wright<sup>1</sup> estimate the rate of increase as follows --

<sup>1</sup> *Diseases of Children*, 1892

At term,	about 2 fluid ounces
Fourth week,	„ 3.4 „
Three months,	„ 5 „
Twelve months,	„ 10 „

The cubic capacity of the skull at term equals 500 c c In the second year it has increased to 1,000 c c In adult life it averages 1,500 c c

The **increase** in the child's **weight** during the first twelve months of its life is very pronounced Pfeiffer gives the following table (abridged) —

	Pounds	Ounces
In the first month,	8	5½
„ third month,	11	15
„ sixth month,	16	3½
„ ninth month,	20	1
„ twelfth month,	22	7

The weight does not increase so rapidly after the end of the first year It is again doubled at the end of the sixth year, and also at the end of the fourteenth year These figures obviously presuppose that the nutrition of the child is progressively maintained

The average length of a child at the end of the fourth year is double the average length at term

At birth, the angle formed by the ramus and the body of the lower jaw is obtuse, being equal to about 140° The body is almost semi-circular in form,

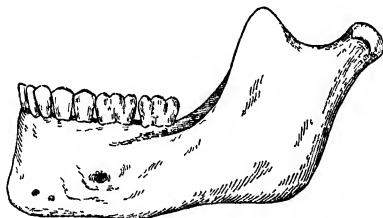


Fig 1 —The lower jaw at puberty

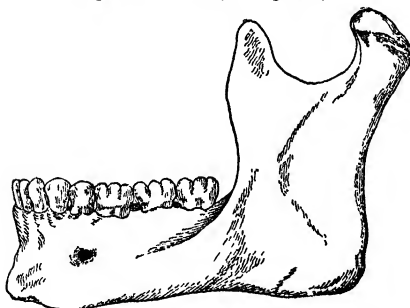


Fig 2 —The lower jaw of adult age

it is shallow, and chiefly consists of the alveolar portion, the basal part is but little developed The permanent teeth being more numerous than the temporary

teeth, require additional space, this is provided by growth of the body of the jaw posteriorly, which changes its form from a semicircle to that of a horse-shoe. Coincident with its increase in length, the body becomes deeper and thicker. The ramus lengthens, and the angle formed by it with the body becomes less obtuse, so that in adult life it approaches a right angle. In advanced old age the teeth are lost, and the alveolar portion of the jaw atrophies, consequently the body is again shallow, and the angle formed by it with the ramus

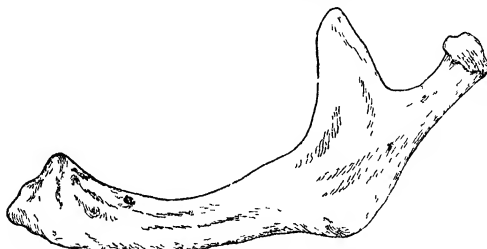


Fig 3—The lower jaw of old age

is once more obtuse. It will be seen from this description that the shallow jaw of infancy and that of old age are anatomically complementary the one to the other—the body of the infantile jaw is almost exclusively alveolar, that of old age is exclusively basal. This is demonstrated by the position of the mental foramen in the infantile and senile jaw respectively—in the former the foramen is low down—in the latter, it opens at the upper border.

### THE TEETH.

The **temporary** or milk teeth are twenty in number. They appear in the following order—

7th month,	lower central incisors
8th „	upper central incisors
7th to 9th „	upper lateral incisors
10th to 12th „	lower lateral incisors
14th „	first temporary molars
17th „	canines
22nd to 24th „	secondary temporary molars

In weakly children, especially in those suffering from rickets dentition is retarded. The converse of this occurs at rare intervals, children have been born with the incisors cut.

The **permanent** teeth are thirty-two in number. They appear in the following order—

7th year,	first molars	
8th „	central incisors	
9th „	lateral incisors	
10th „	anterior premolars, or bicusps	} these replace the temporary molars.
11th to 15th „	posterior premolars	
11th to 13th „	canines	
13th to 16th „	second molars	
18th to 30th „	third molars, or wisdom teeth	



TABLE SHOWING THE PERIODS AT WHICH POINTS OF OSSIFICATION APPEAR AFTER BIRTH

YEARS OF LIFE	BONES IN WHICH THE OSSIFIC POINTS APPEAR
1st	Fourth piece of the body of the sternum
"	Coracoid process of the scapula
"	Head of the humerus
"	Os magnum (carpus)
"	Head of femur
"	Upper end of tibia
"	External cuneiform (tarsus)
2nd	Lower end of radius
"	Unciform (carpus)
"	Lower end of tibia
"	Lower end of fibula
3rd	Great tuberosity of humerus
"	Patella.
"	Internal cuneiform (tarsus)
3rd to 4th	Upper end of fibula
4th	Great trochanter (femur)
"	Middle cuneiform (tarsus)
4th to 5th	Scaphoid (tarsus)
"	Lower end of ulna
5th	Lesser tuberosity (humerus)
"	Internal condyle (humerus)
"	Trapezium and semilunar (carpus)
5th to 6th	Upper end of radius
6th	Scaphoid (carpus)
7th	Trapezoid (carpus)
10th	Upper end of ulna
12th	Pisiform (carpus)
13th to 14th	External condyle (humerus)
"	Small trochanter (femur)

TABLE SHOWING THE PERIODS OF UNION OF EPIPHYSES WITH SHAFTS OF BONES, AND OF BONES WITH EACH OTHER

YEARS OF LIFE	EPIPHYSES AND BONES
1st or 2nd	Symphysis of lower jaw
2nd	Frontal suture, unites from below upwards, may persist
"	Anterior fontanelle filled up
7th or 8th	Rami of ischium and pubis
17th	Epiphysis of upper end of ulna
"	small trochanter (femur)
"	condyles (humerus)
"	upper end of radius
18th	great trochanter (femur)
"	lower end of tibia
"	Lower sacral vertebrae.
"	Portions of acetabulum united
19th	Epiphysis of head of the femur
20th	" humerus
"	lower end of radius
"	" ulna
21st	upper end of tibia
"	lower end of fibula
24th	upper end of fibula
25th	Second and third pieces of sternum
"	First and second sacral vertebrae
"	Epiphysis of clavicle
"	lower end of femur
40th	Manubrium with body of sternum



## CHAPTER VI

### MODES OF DYING.

It is customary and convenient to speak of death as beginning in one of the three essential organs concerned in the maintenance of life—the brain, the lungs, and the heart, failure of action on the part of any one of these organs speedily interferes with the function of the other two. If the blood is insufficiently aerated in the lungs, the vasomotor centres are irritated, and consequently the heart's action is impeded by the narrowing of the blood-channels, the musculature of the heart itself being enfeebled by impure blood-supply. If the heart does not propel the blood with sufficient activity through the lungs, the respiratory centres are ultimately paralysed. Again, if a blood-clot presses on the centres in the medulla, both heart and lungs succumb. If the final obvious indications of life are to be accepted, the heart and the lungs are the organs which by cessation of function actually bring about somatic death. From this aspect the mention of death beginning in the head may be regarded as unnecessary, it is convenient, however, to retain Bichat's classification. It is to be borne in mind that, chiefly, the medical jurist is only concerned in the investigation of deaths which have resulted from violence, but not unfrequently he is called upon to investigate cases in which death was the result of disease—the manner of death, or the circumstances under which it took place, being suspicious of foul play.

The three modes of dying are—*Asphyxia, Syncope, Coma*

### ASPHYXIA.

When the respiratory function is arrested beyond a certain limit asphyxia is the result. There are various ways in which the interchange which normally takes place in the lungs between the blood and the air may be interrupted, viz --

The nervous supply to the respiratory muscles may be abolished either centrally (medulla) or peripherally (pneumogastrics or phrenics). Fixation of the respiratory muscles (tetanus or strychnine), mechanical pressure on thorax, collapse of lungs (pneumothorax), foreign bodies in the air-passages, or closure of them by external compression (strangulation), drowning, respiration deficient in oxygen, spasm of glottis from mechanical irritation (particles of food), or from irritant gases (Cl, SO<sub>2</sub>) are each capable of producing death from asphyxia.

**Symptoms.**—The phenomena of asphyxia may be divided into **three stages**. In the **first**, the respirations are deeper, more frequent, and more laboured than in the normal condition. The extraordinary muscles of respiration are called into action, and the struggle for air becomes more and more severe. The blood becomes more venous, and stimulates the respiratory centres, evoking violent attempts at respiration. In the **second stage**, the inspiratory muscles are less active, while the expiratory muscles contract energetically, as do also almost all the muscles of the body, producing general convulsions. In the

**third stage**, the respiratory centres are paralysed. The pupils are widely dilated, consciousness is abolished and the reflexes are absent. A few gasps at long intervals, and all is over. Hughlings-Jackson<sup>1</sup> directs attention to absence of the knee-jerk when the blood is highly venous, in the earlier stages of asphyxia the knee-jerks are exaggerated, but when the third stage is reached they are entirely lost.

**Post-mortem Appearances.** The right side of the heart, the pulmonary artery, the venæ cavæ, and the veins of the neck are gorged with dark venous blood. The left side is comparatively empty from post-mortem contraction (see under "**Cadaveric Rigidity**"). The blood, nearly black, contains a large amount of  $O_2$ , and therefore coagulates slowly. The hæmoglobin is almost entirely reduced, ordinary venous blood contains a considerable amount of oxyhæmoglobin as well as reduced hæmoglobin (Landois and Stirling).

### SYNCOPE.

When the circulation suddenly fails, syncope is the result. The circulation may fail from cessation of the heart's action (the result of disease (aortic regurgitation, fatty heart etc.), of inhibition (psychical shock, blow on the head, or reflexly from blow on epigastrium). The circulation may also fail from loss of blood (wounds of the large blood-vessels, or of the heart itself, profuse hæmatemesis, etc.) or from sudden withdrawal of blood from the circulation without loss (blows on the abdomen by paralyzing the splanchnics may enlarge the vascular area of the abdomen to such an extent as to deplete the rest of the system).

**Symptoms.** Pallor of the face including the lips, dimness of vision, cold clammy sweat, sense of impending dissolution, craving for more air, great restlessness, gasping for breath, nausea, possibly vomiting, rushing sounds in the ears, momentary delirium quickly passing on to insensibility, followed by convulsions, precede death. The whole of these symptoms are not always present. In simple fainting there may be only immediate loss of consciousness with cold surface and sighing respiration. In all cases the pulse is weak, irregular, or imperceptible. The condition called collapse, though attended by failure of the heart's action, differs from syncope, inasmuch as the patient retains consciousness.

**Post-mortem Appearances.** When death has resulted from insufficient supply of blood to the heart, that organ has been found contracted and empty. When the cause of death has been heart-paralysis, both sides have been found to contain blood. (See "**Cadaveric Rigidity**").

### COMA.

When from any cause affecting the brain insensibility is produced which terminates in death, the individual is said to die from coma. Increase of intracranial pressure, or dynamic disturbance of the cerebrum or of its circulation, may produce coma (concussion, hæmorrhage, tumour, abscess, embolism, thrombosis, depressed fracture of the skull), inflammatory processes (meningitis, etc.), abnormal condition of the blood circulating through the brain (uræmia, certain poisons, as opium, alcohol, and that which produces the complication attending diabetes, known as diabetic coma).

<sup>1</sup> *British Med Journ*, 1892

**Symptoms.**—The symptoms produced by many of the causes of coma above enumerated may take the initial form of stupor, from which the patient may be partially roused for a few seconds or more, this condition subsequently deepens into profound insensibility, from which the patient cannot be roused. Some of the causes enumerated produce sudden coma without any antecedent stupor. In stupor the reflexes may be retained or even exaggerated, in coma they are usually diminished or lost. Power to swallow fluids is consistent with stupor, but not with coma. A comatose person is utterly insensible to all external impressions—he lies powerless, breathing heavily, with stertor from paralysis of the soft palate. The surface is usually covered with a cold sweat, the temperature being at or below normal, except in lesions of the pons and a few other conditions. The pulse may vary, but is often full and laboured. The breathing becomes more and more embarrassed from diminished activity of the respiratory centres, and mucus collects in the air-passages, causing the form of breathing known as ‘the death rattle.’ The pupils, either dilated or contracted, are insensitive to light, and the conjunctival reflex is lost.

**Post-mortem Appearances.** In some of the conditions which produce coma, examination of the brain reveals the cause. From what has been already said it will be apparent that the condition of heart and lungs is not constant, as a rule it resembles more or less that found in death from asphyxia.

### SUDDEN OR UNEXPECTED DEATH FROM NATURAL CAUSES.

Death from natural causes occurring suddenly, or unexpectedly, especially if under suspicious circumstances, is not unfrequently the subject of medico-legal investigation. It is convenient to classify deaths of this kind under three heads. (1) Deaths due to a disease which is universally recognised as being liable suddenly to terminate fatally, and which, as a rule, leaves satisfactory post-mortem evidence of its presence. (2) Deaths due to a disease which, when fatal, does not usually end life abruptly, and which consequently is not generally regarded as a cause of sudden death, in such cases post-mortem examination frequently yields inferential rather than conclusive indications of the cause of death. (3) Cases in which no ascertainable disease is present and consequently post-mortem examination yields absolutely negative results.

1. About one-half of the sudden deaths from natural causes in adults are due to some form of heart disease which has existed for a considerable time, comprising—valvular disease, disease of the coronary arteries, fatty heart, and sclerosis from chronic myocarditis. Spontaneous rupture of the heart, mostly in men, may occur, the seat of rupture is almost invariably towards the front of the left ventricle, in traumatic rupture of the heart the right side, usually the auricle, suffers more frequently than the left in the proportion about as 70 is to 54.

Congenital heart disease is a frequent cause of sudden and unexpected death in infants, and has been found in many cases where, owing to the lividity of the skin and other signs of asphyxia, death from overlying had been suspected. A comparatively small interference in the access of fresh air may prove fatal to an infant whose circulation is already embarrassed. According to Spilsbury,<sup>1</sup> a widely patent ductus arteriosus is the commonest defect, and a patent foramen ovale is fairly common.

Apoplexy and other cognate brain lesions rank second as causes of sudden

<sup>1</sup> ‘Sudden Death,’ *Practitioner*, 1917

death, it is to be borne in mind that small aneurisms of the cerebral vessels occasionally occur in young people, even in children, and by bursting spontaneously cause sudden death from apoplexy. Chronic alcoholism, a potent factor among the causes of sudden death, is frequently associated with rapidly terminating heart and brain disease. Sudden death from asphyxia may be due to œdema of the glottis, membranous deposit in the trachea, pressure of a neoplasm on the trachea, spasm of the vocal cords, whooping-cough, asthma, pulmonary embolism, air embolism, rupture of a vessel or of an aneurism into the air-passages, pneumo- and hæmo-thorax, pleuritic effusion, and epilepsy. The rupture of a gastric or of an intestinal ulcer, of an aneurism, of a varicose vein, of the surroundings of an ectopic gestation, the formation of a peri-uterine hæmatocele, may severally prove quickly fatal. Nephritis (uræmia and apoplexy), diabetes, exophthalmic goitre, and Addison's disease may also terminate with unexpected rapidity. Hæmorrhage into the pancreas occasionally causes sudden death, apparently from the impression produced on the contiguous nerve-centres, it is most common in males over forty years of age, but it has occurred in a woman aged twenty-four, heart disease, obesity, and the abuse of alcohol appear to be predisposing causes. It sometimes occurs, however, in spare abstemious people free from obvious disease. In several instances sudden death has followed spontaneous rupture of an enlarged spleen, the result of tropical malarial influences, the individuals immediately before rupture being apparently quite well. A mere pat with the palm of the hand might rupture such a spleen, and death thus caused might be attributed to criminal violence. The most careful post-mortem examination may fail to reveal the cause of death in some of these diseases— as spasm of the vocal cords, whooping-cough, and asthma. Epilepsy may or may not leave characteristic appearances, such as recent injury to the tongue, ecchymoses on the conjunctivæ, the face, the neck, and occasionally on the trunk.

2. Certain diseases which, when fatal, do not usually end suddenly may exceptionally do so. Such are — Abscesses and tumours of the brain, meningitis, diphtheria, tracheitis, enlarged thyroid, pulmonary apoplexy, pneumonia, phthisis, pneumothorax, pleurisy, ulcerative endocarditis, old adherent pericardium, pericarditis, volvulus and other causes of acute intestinal obstruction, peritonitis, gall-stone colic, rupture of an echinococcus of the liver or elsewhere, influenza, gout, acute rheumatism, and Bright's disease. When fully developed many of these diseases are usually inhibitive of locomotion, but occasionally they occur in a latent or an ambulatory form and terminate suddenly without their presence being even suspected. Öllivier<sup>1</sup> records the case of a man who was taken ill whilst at work, he went to bed and died in four hours, and at the autopsy the cause of death was found to be purulent meningitis. In children, especially, death may unexpectedly occur from septic meningitis due to long-standing middle-ear disease. The advance of an acute disease towards a fatal issue may be excessively rapid. Strassmann<sup>2</sup> examined the bodies of two young women who went to bed not feeling quite well, although they had done their usual day's work, the following morning they were discovered dead in bed, the cause of death being a diphtheritic deposit which had travelled rapidly down the bronchi. Diphtheria may cause unexpected death, during the stage of convalescence, from heart failure, which may occur after a *latent attack of diphtheria that has escaped notice*. Guthrie<sup>3</sup> saw a girl aged two years with diphtheritic

<sup>1</sup> *Arch. gen. de Méd.*, T. 1, Serie 3e

<sup>2</sup> *Lehrbuch d. gericht. Med.*, 1895

<sup>3</sup> *The Lancet* 1894

vulvitis who died very suddenly from cardiac failure, in many instances such an attack and even one affecting the pharynx may pass unobserved. After diphtheria a clot may form in the right ventricle and may cause sudden death from embolism.

When an enlarged thyroid causes sudden death the gland is usually enormously hypertrophied, sometimes hæmorrhage takes place into a cyst and determines rapid suffocation from increase of pressure which previously was all but sufficient to provoke fatal dyspnoea. Exceptional causes of death from suffocation may occur, a man in his usual health died quite suddenly, and at the autopsy a round worm, six inches long, was found in the left bronchus, the characteristic signs of death from suffocation being present. The case narrated on p 177 affords another illustration. The contents of the stomach may be vomited and a portion of them may be drawn into the air-passages and thus cause sudden death, for examples see p 176.

Pleurisy with effusion may exist (without materially interfering with the breathing) in a man going about as usual who, after slight exertion or trifling violence or even in the absence of both, may die instantaneously. Brouardel<sup>1</sup> relates the case of a policeman who, on receiving a blow on the chest from the fist of a man whom he was arresting, fell down dead, the right pleural cavity was found full of fluid, and no other indication of the cause of death was present. In pleurisy with effusion there may be great risk of sudden death on mere change of posture—as in getting out of bed—and also from the passage of an aspirating needle into the pleural cavity, death occurring either immediately or after an interval of several hours, without any post-mortem appearances beyond those usually produced by the disease when it has run its ordinary course. Pneumonia may be latent and may cause death exceedingly abruptly. Strassmann<sup>2</sup> relates the case of a man aged fifty who, whilst inquiring for work at a registry office, fell down dead, and at the autopsy the whole of the upper lobe of the right lung was found in a state of grey hepatisation. In old people latent pneumonia is a not unfrequent cause of sudden death, the individual apparently being in his usual health up to the time of his death, post-mortem examination reveals nothing beyond the anatomical changes common to ordinary fatal cases of this disease. An overloaded stomach may determine speedy death in a patient suffering from a disease which occasionally terminates suddenly. Sturges and Wilkins<sup>3</sup> recorded the case of a boy aged seven, apparently in good health, who ate a hearty dinner, and immediately after played cricket for an hour and a half, he then ran to school, fell down insensible, and died forthwith, at the autopsy recent pericarditis was found to be the sole lesion. In a large number of cases of sudden death the victim was undergoing some form of physical exertion at the time, such as playing a fatiguing game, hastening to catch a train, mounting stairs, defæcating, or was engaged in sexual intercourse.

A case recorded by Rake<sup>4</sup> illustrates the occurrence of sudden death from acute intestinal obstruction, a strong man aged fifty, whilst gardening, was suddenly attacked with vomiting and abdominal pain, he died in two hours, and at the autopsy a volvulus of the transverse colon was found, death being probably due to disturbance of the solar plexus. Occasionally rupture of an intestinal ulcer occurs in cases of latent or ambulatory enterica, sudden death also occurs in this disease apart from rupture of an ulcer, it most frequently occurs about the beginning of the third week, and is apparently due to rapid

<sup>1</sup> *La mort et la mort subite*, 1895

<sup>2</sup> *Lehrbuch d. gerichtl. Med.*, 1895

<sup>3</sup> *The Lancet*, 1885

<sup>4</sup> *The Lancet*, 1889

or instantaneous heart failure Dewevre<sup>1</sup> collected statistics which indicate that sudden death from typhoid fever reaches its maximum between the ages of twenty and twenty-five years, and that it is commoner in men than in women. In children, sudden death may occur in the early stage of enterica just as the intestinal follicles are beginning to swell.

Bright's disease, both acute and chronic, occasionally causes sudden death apart from uræmia, or from apoplexy, in advanced contracting kidney, sudden death may occur without any post-mortem indications of the ultimate cause of death either in the brain or elsewhere. Occasionally sudden death occurs to elderly people afflicted with gout, and also to those who are excessively corpulent, in some instances without affording any post-mortem evidence as to the actual cause of death. Kisch<sup>2</sup> records nineteen cases of sudden death in obese persons, twelve from acute œdema of the lungs, six from cerebral hæmorrhage and one from rupture of the heart, seven were under and eleven were over fifty years of age, fifteen were men and four were women. Apart from cases of hyperpyrexia, acute rheumatism may cause unexpected and sudden death.

In most deaths under anæsthetics, the morbid condition which has necessitated the administration of the anæsthetic has been mainly responsible for the fatality. Malignant growths have figured prominently in these deaths and often the patient has been so ill that the anæsthetic can only have played a small part in bringing about the death. In a number of instances, however, of such deaths, where neither the operation nor the morbid condition appeared to be so severe as to be the cause, the condition known as *status lymphaticus* has been found on post-mortem examination. The disorder has usually been unsuspected during life. *Status lymphaticus* is characterised by persistence of the thymus and an increase in the lymphoid tissue in various parts of the body. The lymphatic glands and the spleen are enlarged, and the heart shows degenerative changes, usually of a fatty character. Spilsbury,<sup>3</sup> in an analysis of 182 deaths occurring under anæsthetics, noted *status lymphaticus* in 48 cases. He found a much greater proportion to occur under chloroform than under other general anæsthetics, and he considers the decrease in the number of deaths under anæsthetics during recent years to have been due to the extended employment of open ether in place of chloroform. A condition of dyspnoea due to pressure of an enlarged thymus has been described by various writers, but, according to Thursfield,<sup>4</sup> the cases are extremely rare. He considers that the enlargement of the deep-seated glands is more likely to be the cause of the pressure than the enlarged thymus, and brings forward reasons for supposing that in many cases some factor other than pressure is the cause of death.

3 When death suddenly occurs without affording any post-mortem indication it is often determined by an external impulse acting either (a) directly, or (b) mediately on the nervous system.

(a) Several cases are on record in which an individual, whilst in a state of extreme mental excitement, has died suddenly solely from psychical stimulation. Marchal<sup>5</sup> relates that a military prisoner was transferred to the hospital, and whilst the doctor was reading the statement of his previous behaviour, the prisoner, who had a profound dread of being sent back to prison, was observed to be greatly perturbed, and he suddenly fell back dead, at the autopsy no lesion other than engorgement of the cerebral meninges was found. See also the cases quoted from Maschka and Templeman on p 254. Death occurring as

<sup>1</sup> *Arch. gen. de Med.*, 1887.

<sup>2</sup> *Berliner klin. Wochenschr.*, 1886.

<sup>3</sup> *Op. cit.*

<sup>4</sup> *Transactions of the Royal Society of Medicine*.

<sup>5</sup> *Bonn, De la mort subite*.

in these cases is usually attributed to "heart disease", a distinction, however, is to be drawn between death due to disease of the heart and death due to profound perturbation of the nerve-centres causing inhibition in a heart in which there is no trace of disease. When searching after death for evidence of disease in a suspected heart it is to be borne in mind that absence of atheroma at, or of narrowing of, the mouths of the coronary arteries is not sufficient to exclude angina as a cause of death, it is necessary to lay open the entire length of the coronary arteries, since the seat of obstruction may be some distance from the orifice, which itself may be quite patent.

(b) When the external impulse acts mediately on the nervous system it is by means of some slight physical impact, or injury, that in itself is a totally inadequate cause of death, but which occurring during a period of acute apprehension, or of extreme terror, may suddenly and completely inhibit, or in some instances profoundly depress the heart's action, in the latter case death may not take place for several hours after the occurrence of the causal event, as in the following case related by Beaunis<sup>1</sup> - A man whilst shooting wild beasts was pounced on by a panther, some Arabs who were present immediately shot the animal, and on examining the victim found that the panther's claws had only inflicted a few pricks on the shoulder, so slight that they scarcely bled, the man remained in a state of profound terror, and died on the following day. At the autopsy no lesion was found that would account for death the claw-marks were quite superficial. A still more remarkable case which occurred in India is recorded by Francis<sup>2</sup>. A man was roused out of sleep by feeling something creeping over his naked legs, having an intense dread of reptiles, and imagining that it was a cobra, he became collapsed and died in six hours, although before death took place it was discovered that a harmless lizard had been the cause of his fright, the patient, however, was too profoundly collapsed to recover. The immediate lethal effect literally of a mere touch is illustrated by an instance which occurred to Dupuytren,<sup>3</sup> who, when about to perform lithotomy on a boy, demonstrated to his students the line of the projected incision by drawing his finger-nail along the perineum, the boy immediately fell back dead. A surgeon was examining, with a catheter, the bladder of a robust woman, she closed her eyes and died forthwith. Bonvalot<sup>4</sup> relates that as the canula of a syringe was introduced into the os uteri of a young woman with the object of procuring criminal abortion she fell back before any of the fluid had been injected, and died in a few minutes. Cases are on record in which sudden death occurred during the actual administration of an intra-uterine injection. Spilsbury<sup>5</sup> gives the following description of a case he saw - -

A young married woman, who had had one child, was left alone in her house one evening for about 1½ hours. Her husband on his return found her in her night attire lying on her back on the bedroom floor dead. On a sheet spread out on the floor beneath her was an enema syringe, one end of which was in a pudding basin half filled with a concentrated warm soap solution which set solid on cooling. Similar soap solution was found in the bulb of the syringe. The other end of the syringe was blood stained and lay between her legs. On post-mortem examination, she was a healthy woman about 2½ months' pregnant. The neck of the uterus was slightly dilated, and there were several small lacerations around the external os. The uterus contained a complete ovum with the membranes unruptured, but separated from the wall for some distance upon the posterior surface, the space being occupied by about 1½ ozs. of soap solution. No serious injury had been inflicted, and she

<sup>1</sup> Vincent, *Des causes de la mort prompt*

<sup>2</sup> *Med. Press and Circ*, 1883.

<sup>3</sup> Nussbaum, *Ueber Unglücke in der Chirurgie*

<sup>4</sup> *Annales d'Hygiène*, 1892.

<sup>5</sup> *Op cit*

had died suddenly from the shock of injection of fluid into the uterus for the purpose of procuring abortion. The evidence failed to show that any other person had assisted her in the commission of the act.

In two very similar cases, mentioned by the same physician, there was slight dilatation of the external os uteri, but no fluid had been injected into the uterus.

On more than one occasion women have died suddenly at the moment a simple digital vaginal examination was being made. Brouardel<sup>1</sup> saw one such case in hospital, and states that he knows of two others which occurred in the consulting-rooms of medical men. It is to be borne in mind that the bladder, urethra, and genitals are comprised among those parts of the body, the abrupt handling, or compression, of which is attended with exceptional risk of sudden death from cardiac inhibition, the other parts are—the nasal passages, the larynx, and the epigastrium. (See article by the author on "Sudden Death," *The Lancet*, 1897.)

## CHAPTER VII

### SIGNS OF DEATH, AND CHANGES IN THE BODY AFTER DEATH.

THE movements which accompany respiration and circulation are the two most obvious indications of the existence of animal life. If signs of movement of the lungs and of the heart are present, it is clear that death has not yet occurred. If the lungs cease acting, the heart may continue to beat for many minutes, which, also, is conclusive that death has not yet taken place. When both heart and lungs have ceased to act, the tissues retain 'vitality' for some time, during the continuance of which it cannot be truly asserted that the individual is dead. To this lingering "vitality" of components is due the potentiality of resuscitation of the entity they compose, after temporary cessation of the main functions by which life is sustained. If the heart persistently ceases to beat, the vitality of the tissues progressively diminishes, until at last a point is reached when death is absolute and universal, when this point is reached, but not before, resuscitation is impossible.

It is very necessary for the medical jurist thoroughly to appreciate the importance of the distinction between *systemic* or *somatic death*, and *death of the tissues* or *molecular death*. In profound syncope for example, there is every external appearance of death, but, if the heart resumes its function, the patient recovers. Hence the necessity of careful study of the phenomena which succeed death, in order to be able to distinguish between real and apparent death.

Auscultation, carefully conducted and repeated if necessary at intervals, will enable an opinion to be formed as to whether the heart and lungs have or have not ceased to act. Absolute silence should be maintained during the investigation, which should be conducted with all deliberation. Error on the part of the practitioner, should he wrongly pronounce that death has taken place, is so obviously capable of refutation that his reputation is damaged, and what is of infinitely greater importance, such an error might lead to that

<sup>1</sup> *La morte et la morte subite*, 1895



most ghastly of all blunders—the treatment of a living being as though he were dead. A number of tests have been devised in order to determine whether the circulation has or has not ceased, such as examining the hands of the supposed dead person by transmitted light, during life the margins, the tips, and the web of the fingers appear red and translucent, after death the fingers appear opaque up to their edges. On tying a piece of string round a finger its distal end becomes swollen and bluish-red if the circulation continues, if it has ceased no effect is produced. Icard<sup>1</sup> proposed to inject subcutaneously a solution composed of one gramme each of fluorescin (resorcin-phthalein) and of sodium bicarbonate dissolved in eight cubic centimetres of water, if the circulation continues, the entire skin and mucous membranes in a few minutes assume a yellowish-green colour, and fluorescin may be detected in the blood by passing a few threads of cotton under the skin (at a part distant from the seat of injection) and, after their withdrawal, by boiling them in a little water a green coloured solution is obtained, if the circulation has ceased no such effects are produced.

After the occurrence of somatic death—that is, after permanent cessation of respiration and circulation—certain phenomena occur in definite order. The time occupied by many of these phenomena, although variable, is limited, and, consequently, by ascertaining the stage at which they have arrived, an opportunity is afforded of approximately estimating the interval that has elapsed between death and the period of investigation. These phenomena are manifestations of the occurrence of molecular death. The rapidity of their onset is, to some extent, determined by conditions which exercise an influence on the amount of vital energy with which the tissues are endowed immediately before somatic death. Such conditions comprise disease, poison, violent exercise, and other exhausting influences. If the tissues are depressed in vitality at the moment of somatic death, they rapidly succumb to molecular death, and the course of the succeeding phenomena is hastened. If, on the other hand, they are in full activity when somatic death takes place, they resist the advent of processes which are really the antecedents of decomposition.

### CESSATION OF RESPIRATION AND CIRCULATION.

At the moment of somatic death, the skeletal muscles lose their tonus and become flaccid. When death takes place whilst the body is in the recumbent posture, the loss of tonus is immediately manifested by dropping of the lower jaw, the eyelids remain open, or partially so, and the limbs are flexible. The pinched look of the face—*facies hippocratica*—that frequently manifests itself at the time of death, changes to a more or less peaceful expression, often—in the case of persons above middle age not much emaciated—evoking the comment from the bystanders that the deceased now resembles the appearance he presented in youth, this alteration in facial expression is due to relaxation of the muscles smoothing away the sharp contour of the features. The eyes lose their lustre and acquire the peculiar ghastly stare so characteristic of death. The pupils dilate at the time of death, and do not respond to light. Marshall<sup>2</sup> states that contraction of the pupils subsequently takes place for a period varying from one to forty-eight hours after death, that atropine dilates the pupils after death, in some instances as long as four hours after, and that eserine contracts the pupils, but not for so long after death. The surface of the body, including the lips, becomes pallid, people of florid complexion often retain

<sup>1</sup> *La Mort réelle et la Mort apparente*, 1897.

<sup>2</sup> *The Lancet*, 1885.

their colour for several days after death. Ten or twelve hours after death, the eyeballs sink in the orbits and become flaccid, so that the cornea retains the dint caused by pressure of the finger-nail or other hard substance.

### POST-MORTEM COOLING.

Death abolishes the distinction between living and inanimate matter, and consequently the human body after death becomes subject to the laws which govern inanimate matter. That remarkable property with which man in common with other animals is endowed—the power of maintaining an equable internal temperature, whatever may be the temperature of the surrounding air—is now lost, the consequence is that the body yields up its heat and slowly but progressively cools down to the temperature of the surrounding media.

From **fifteen to twenty hours** is the time usually required for the body to cool to the temperature of its surroundings. It is obvious that this can only be accepted as a general statement. The actual time taken by a cadaver to become cold is determined by conditions, some of which are of **internal** and others of **external** relation.

**Conditions of Internal Relation.**—At the time of death resulting from some diseases, and from certain other modes of death, the body-heat greatly exceeds the normal. This has been observed in cholera, acute rheumatism, poisoning by strychnine, tetanus, and in some forms of apoplexy. After death from cholera, yellow fever, and some other diseases, post-mortem elevation of temperature has been observed. Under like external conditions, a cadaver having at the moment of death a temperature of 8° or 10° F. above normal, would take longer to cool than one at the normal temperature. Certain modes of death, without necessarily causing any elevation of temperature, retard the rate of cooling: death from suffocation, and some other kinds of sudden death from violence, may have this effect. The state of nutrition at the time of death also exercises a considerable influence; bodies loaded with fat do not cool so rapidly as those which are emaciated. The bodies of persons in the prime of life do not cool so rapidly as those of very young or of very old people. Those who have died from lingering or wasting diseases cool rapidly, the temperature probably being subnormal before death.

**Conditions of External Relations.**—The temperature of the surrounding media and their capacity for heat-conduction exercise an important influence; a body submerged in water will cool more rapidly than in air, because water is colder and is a better conductor of heat than the air above it; a body lying naked on the flags will cool quicker than one protected by clothing and lying on a bed. Without particularising further, it may be accepted that the greater the difference of temperature between the dead body and the surrounding media, the more rapidly is its heat lost in a given unit of time; therefore, the rate of post-mortem cooling is not uniform, but is proportional to the difference in temperature existing at any given period between the body and its surroundings. From this it follows that the cooling rate will be quicker in the earlier than in the later hours, because in the later hours the temperature of the body more nearly approximates to that of its surroundings; during the first few hours it may average 2° or 3° F. per hour, subsequently it will not exceed 1° F. per hour. The longer of the two periods—twenty hours—stated as sufficient in ordinary cases to allow the body to cool, may be accepted as practically, but not absolutely, correct. For the reason just given, the cooling-rate is exceedingly slow when the temperature of the body differs but slightly

from that of its surroundings, therefore, the last few degrees of body-heat take a long time to disappear, the body in the meantime being cold for all practical purposes. It is to be borne in mind that the loss of body-heat must be *progressively continuous* before it is accepted as a sign of death, the temperature at any given moment is not trustworthy, since it may be considerably subnormal during life.

### POST-MORTEM STAINS.

Coincident with cooling is the formation of certain discolorations on the dependent parts of the body. Various terms are used to indicate these stains, as—cadaveric hypostases, cadaveric ecchymoses, post-mortem lividities, sugillations. When the subject of giving evidence was dealt with, the necessity of avoiding technical terms in the witness-box was emphasised, in furtherance of this advice it is recommended that the words which head this section should invariably be used when discussing the subject, as conveying to the ordinary hearer a clearer idea of what is meant than any of the synonyms. Post-mortem stains are produced by gravitation of the still fluid blood into the capillaries and the venous radicles of the lower parts of the body, and are obviously determined by the position in which it rests. If the body is lying on its back the lobes of the ear, the shoulders, the lumbar region, the buttocks, and the posterior parts of the legs and arms will constitute the lower parts. If the body is in the prone posture, the face, chest, abdomen, and the anterior parts of the legs and arms will be the lower parts. In addition to the results produced by gravity are those due to loss of tonus of the walls of the capillaries which occurs at the time of death. The effects produced by the gravitation of the fluid blood are not limited to the surface of the body, the internal organs are also subject to them.

As the formation of post-mortem stains is dependent on the fluidity of the blood, the question arises—how long after death does the blood remain fluid within the body? Coagulation within the dead body commences later, and takes place much more slowly, than is the case with blood withdrawn from the living organism. It is impossible to state with exactitude the time that elapses between death and the commencement of coagulation, about four hours is the period allotted. Clots may form within the large veins at this interval after death, but the progress of coagulation is so slow—especially in the smaller veins—that the great bulk of the blood is probably still fluid for many subsequent hours. This being the case, the exact period after death when the blood begins to coagulate is of little importance. The time occupied in coagulation may be further prolonged by the condition of the blood at the time of death. The presence in it of excess of carbon dioxide retards coagulation, therefore, the blood of persons who have died from suffocation coagulates very slowly.

The salient features of slow coagulation outside the body are distinguishable in the blood-clots formed within the cadaver. These features are sinking of the red blood-corpuscles to the lower stratum before solidification takes place, and the consequent formation of a more or less colourless layer at the upper part of the clot, this division of the clot into two layers is of importance to the medical jurist. On examination of a dead body in which coagulation has taken place, the clots ought to correspond with the position of the body, the coloured layer being furthest from the upper surface of the body. If this is not the case—if the *colourless* part of the clot is found at the *undermost* part of the body—proof is afforded that the position of the body has been reversed after coagulation.

took place. The blood-clot formed within the body after death is not nearly so firm as one formed from blood withdrawn from the circulation during life. Apart from death from suffocation, it does not appear that the reluctance of the blood to coagulate within the vessels depends upon the presence of  $\text{CO}_2$ , as blood which has remained fluid within the body for some hours after death will coagulate when received over mercury without exposure to air.

To return to post-mortem stains. They begin to appear on the undermost parts of the body from **four to twelve hours** after death, not unfrequently they may be seen earlier than the first-mentioned period. They consist of patches of a dull red, or they may be of a bluish slate colour. At first they impart a mottled appearance to the skin, later on the individual patches coalesce and form large areas of discoloration. The outline of the patches is irregular but is well-defined, the margins stand out in bold relief against the neighbouring uncoloured skin. The stained portions are not elevated above the level of the surrounding skin. Although post-mortem stains occur on the undermost parts of the cadaver, they are absent from those parts which are in contact with the substance on which the body lies. If a body lies in the ordinary position—on its back—the shoulders, buttocks, calves, and heels will be unstained. The reason for this is that the compression of the parts named, caused by the body-weight on the one hand and the resistance of the unyielding surface which supports the weight on the other, prevents the gravitation of the blood into the vessels of the skin. An amount of compression much less than that produced by the weight of the body is sufficient to exclude the blood from the part thus compressed, a tight collar or neckerchief may act in this way, and so produce a mark round or partly round the neck that might resemble to some extent the mark left by a cord after death by strangulation. Post-mortem stains continue to form so long as the blood, or a part of the blood, remains uncoagulated. Whilst the blood remains fluid, the post-mortem stains are only permanent provided the position of the body at the time of their formation remains unaltered. If post-mortem stains have made their appearance on the posterior part of a body which lies on its back, and whilst the blood is still fluid the position of the body is reversed, the stains on the back part will disappear and fresh ones will be formed on the now dependent parts in front. Permanency of position in post-mortem stains is only secured by coagulation of the blood, after it has taken place, any alteration in the position of the body produces no effect on them, nor are new ones formed. For this reason, if the position of a body is materially changed after coagulation, it will be found that the post-mortem stains do not correspond with the altered position, and consequently afford evidence of the body having been interfered with. Casper states that they are invariably present after death from hæmorrhage, other authorities have found them absent, or but feebly indicated in such cases, and also when anæmia from disease was the cause of death.

The distinction of post-mortem stains from bruises made during life is of great importance. Mistakes in this relation may be, and have been, the cause of doing serious injustice to innocent persons. The difference is very marked and easy of recognition. In **post-mortem stains** the blood which produces them is still within the blood-vessels—it is contained in the flaccid, dilated venous radicles and capillaries of the superficial layers of the corium. The surface is uninjured. If examined in an oblique direction, or with an oblique light, no trace of disturbance of the cuticle is found. The parts stained are not elevated, they are practically on the same level with the surrounding skin. The stains are always on the most dependent parts of the body, with the exception of such

parts as are subject, or have been subjected, to compression. The margins of the stains are well defined, they do not fade away into surrounding skin. The depth of colour of the patches of stain is uniform, or nearly so. The stains when in patches (such as might, by the ignorant, be mistaken for bruises) rapidly increase in size, mostly in a direction determined by gravity.

In **bruises** made during life, the discoloration of the skin is caused by extravasation of blood in and under the papillæ of the true skin from vessels which have been ruptured by violence. The surface of the skin will generally be found disturbed, the result of impact with the object that produced the bruise. Except in the case of very slight injuries, the bruised part will be found more or less elevated. The position of the mark of a bruise is not determined by gravity, it may be on any part of the body, and instead of having a well-defined irregular outline, it will correspond somewhat in shape with the causal agent, or with that part of it which came in contact with the surface, and the margin will be ill-defined, fading away into the surrounding skin. The colour of a bruise is not uniform, if the injury has existed for a day or two a zone of yellow or green may be seen round the outer parts.

Such are the differences observable on the surface between post-mortem stains and bruises. **The critical distinction is made by incising the part.** In a post-mortem stain no blood escapes except from a few minute points which represent the divided vessels of the corium previously distended with blood. On the other hand, an incision into a bruise reveals the presence of effused blood, either clotted or fluid, below the rete, or, when the bruise has been the result of great violence, still deeper down.

So far for external post-mortem stains. The same conditions that produce the stains on the skin influence the **internal organs**. When the body has been lying on the back, the veins of the pia mater at the posterior part, and the lateral and occipital sinuses, will be found filled with blood. This is even the case after death from hæmorrhage, a fact to be borne in mind lest the presence of so much blood in the highest part of the body (during life) should be regarded as inconsistent with that cause of death. The posterior fourth of each lung is almost invariably filled with blood, here also there is some risk of attributing the condition to pathological causes. The posterior parts of the stomach and of the intestines, especially those in the pelvis, are also stained, the absence of inflammatory exudation, and the fact that on stretching out the bowels the coloration will be found absent at parts lying between the discoloured portions, is sufficient to prevent error of interpretation. The posterior halves of the kidneys are usually gorged. The veins in the pia of the cord often present an appearance not unlike that resulting from meningitis. The heart is free from staining, but fibrinous clots— the so-called cardiac polypi—are common.

### CADAVERIC RIGIDITY.

The flaccidity of the muscles which occurs immediately after death gives place to an opposite condition of extreme hardness. So long as the muscles remain flaccid they retain their irritability, and respond to electrical stimuli in the same way that they do in life, this is due to the persistence of molecular or tissue life after the cessation of somatic life. At a variable period after death, the muscles of the lower jaw begin to stiffen, this is the first indication of the onset of **cadaveric rigidity** or, as it is also called, **rigor mortis**. Under ordinary circumstances, the skeletal muscles begin to stiffen in from **four to ten hours** after death. The stiffening spreads from the muscles of the jaw to those of the

face, neck, and trunk, and lastly to the limbs. It is fully developed in from two to three hours, when the entire body is firm and stiff. The limbs cannot be flexed at their joints without considerable force, and the body when moved behaves as though it was devoid of articulations. This condition lasts for a period varying from a few hours to six or eight days. Twenty-four to forty-eight hours may be regarded as the average duration of cadaveric rigidity. In addition to the hardness, a certain amount of contraction of the muscles takes place during the onset of cadaveric rigidity, the muscles become shorter and thicker, and the limbs, if left to themselves stiffen in a position of partial flexion. Experimentally it is found that a muscle in the act of stiffening will lift a weight, showing that actual contraction takes place. This question will be fully discussed later on. When cadaveric rigidity is present the muscular electrical current is either abolished, or there is a feeble current in the opposite direction to the normal. The muscles have also lost the power of responding to electrical stimuli. The chemical reaction of the living muscle at rest is neutral or slightly alkaline, that of muscle doing work is acid. Muscles in which cadaveric rigidity is present have a strongly acid reaction, due partly to the formation of sarcolactic acid, and partly to acid phosphates, in addition, a large amount of carbon dioxide is set free. The actual cause of the muscles becoming hard and stiff is the coagulation of the muscle-plasma within the sarcolemmas of the muscular fibres (Kuhne). Several cases of ante-natal rigidity are recorded. An eight months' child was in the state of rigor mortis when born, and the rigor had not passed off on the following day.

The rigidity seems to exist in two conditions the first passing on to the second by simple lapse of time. In the first condition, it is supposed that the muscle-plasma is only partially coagulated, being of a jelly-like consistence, in this stage the muscle has not yet lost its irritability. If, whilst the muscle is in the first stage of rigidity, fresh arterial blood is transfused into the blood-vessels, the semi-coagulated material is dissolved, and it is then found that the muscles are still irritable and respond to electrical stimuli (Brown-Sequard).<sup>1</sup> When the second stage is reached the coagulation of the muscle-plasma is complete, and the irritability of the muscles has permanently disappeared. This coincides with the period of death of the muscular elements, and of the tissues of the body generally. When cadaveric rigidity is fully developed, if one of the limbs is forcibly flexed at a joint, the rigidity does not return, the joint remains supple, and the limb in the position in which it is left. In this way *post-mortem* rigidity may be distinguished from *ante-mortem* rigidity as occasionally met with in hysterical or cataleptic subjects. In the latter condition, if the limb is forcibly bent it tends to return to its previous position, and again becomes rigid. In the first stage of cadaveric rigidity, however—when the myosin is only partially coagulated—a certain degree of stiffening again occurs after a joint is forcibly flexed.

**The conditions which hasten the onset of cadaveric rigidity** are those which have an exhausting or depressing influence on the muscles immediately before death, hence, after violent muscular exercise, death is speedily followed by rigidity. It has been observed that the bodies of soldiers killed at the commencement of a battle, before they have undergone much fatigue, do not become rigid so soon as those of their fellow combatants who succumb at a later period, after many hours' arduous fighting. After death from diseases or poisons, which depress or exhaust the system (especially if attended with clonic spasm

<sup>1</sup> See also Heubel on "Die Wiederbelebung des Herzens nach dem Eintritt vollkommener Herzmuskelstarre" Pfüger's *Archiv*, 1889

of the muscles), rigidity comes on early. Animals that have been hunted for some time before death stiffen almost at once, and within a few minutes may be held out by the hind legs perfectly rigid. On the other hand, in speedy death (with exceptions presently to be mentioned) occurring to individuals in vigorous health, the onset of cadaveric rigidity is delayed. Even in those who are in full vigour, if death is immediately preceded by convulsions, rigidity comes on early, because the vital energy of the muscles is lowered. It has been found that the muscle-currents disappear, and the electrical irritability of the muscles is diminished before the development of rigidity. All these facts point to one conclusion—that cadaveric rigidity is a consequence of death of the muscles, that it creeps on as the muscles are dying, and is therefore, partially present before they are dead, but is not fully developed until molecular death has taken place.

It may be asserted as a general proposition that the **sooner** rigidity comes on after death, the **sooner** will it **pass away**. The converse is equally true—the longer it is in appearing, the longer will it last. An unexhausted condition enables the muscles, within limits, to resist disintegration when deprived of further aid from the nutrient supply and the eliminative agencies by which their vigour was maintained during systemic life. A high state of vigour is resistant to change, not only up to the period when change commences, but for some time after. Hence it is, that both the onset and the duration of cadaveric rigidity (other conditions being equal) are dependent on the store of vitality possessed by the muscles at the moment of somatic death.

**Heat-Stiffening.**—The rigidity of a cadaver that is fully under the influence of ordinary cadaveric rigidity may be increased by subjecting the body to a temperature of  $75^{\circ}\text{C}$ . The explanation is that other albuminates present in the muscles besides myosin are thus coagulated. Myosin coagulates (in mammals) at about  $50^{\circ}\text{C}$ , another albuminate coagulates at  $47^{\circ}\text{C}$ , and serum-albumin coagulates at  $73^{\circ}\text{C}$ . If, therefore, either before or after the full development of cadaveric rigidity, the body is subjected to a temperature exceeding  $73^{\circ}\text{C}$ , but short of causing disintegration, all these albuminates are coagulated, and a higher degree of rigidity is produced than that dependent on natural causes.

**The Involuntary Muscles** are also affected by cadaveric rigidity, the only internal organ necessary to consider in this respect is the **heart**. Very soon after death, before the skeletal muscles begin to stiffen, the heart muscle becomes rigid. It has long been observed that when the heart is in the condition of cadaveric rigidity, the ventricular walls, especially the left, are firm and contracted, and present an appearance totally different from that which obtains after rigidity has passed off. It has also been observed by physiologists that certain changes in the condition of the heart-muscle take place during the development of cadaveric rigidity. The heart, like the skeletal muscles, not only stiffens, but it undergoes contraction after death sufficient to alter entirely the relative capacity of its cavities. This fact is of great importance to the medical jurist, since a decision as to the mode of death is not unfrequently based on the condition of the heart as found at the necropsy. When the post-mortem appearances of death from syncope were described, the condition of the heart was represented as being contracted, which is often the case at a necropsy on the body of one who has died from syncope, the result of want of blood to supply the heart. The question arises—is this the condition of the heart at the time of death, or, in other words—did death take place with the heart in systole? Strassmann's<sup>1</sup> experiments are very instructive in relation to this question.

<sup>1</sup> *Vierteljahrsschr f ger Med*, 1889, 1896

In a number of animals which had been killed by sudden paralysis of the heart, he opened the thorax as soon as circulation and respiration had ceased. In some the heart was examined immediately, others, after superficial investigation, were covered over and left until the next day. When the heart was examined immediately after death, the left ventricle was found to contain more blood than the right, but when the examination was made on the following day, the left ventricle was almost invariably found firmly contracted, and its contents for the most part, or entirely, expelled either into the left auricle or into the aorta. In animals that had been killed by rapid suffocation, when the heart was examined immediately after death, the right ventricle contained on the average twice as much blood as the left, at a subsequent examination, the left ventricle was found contracted and empty, or nearly so. In all cases, when the heart was examined immediately after death, even after death from strychnine, both right and left ventricles were found in diastole, relaxed and filled with blood, in no case was the heart found to have stopped in systole. Strassmann directs attention to the extremely early development of cadaveric rigidity in the heart in animals poisoned with hydrocyanic acid it was observable one hour after death, even so early as this, the left ventricle did not contain nearly so much blood as it did immediately after death. Post-mortem contraction of the right ventricle is much feebler than that of the left, only in cases where hæmorrhage has taken place is it found empty. These investigations indicate that the condition of the heart at the necropsy affords no certain proof of its condition at the moment of death, only when left and right sides are both found to be filled with blood can it be truly said that the original condition has been maintained.

In post-mortem examinations on the human subject, it is usual to allow an interval of at least twenty-four hours to elapse before the examination is made. During this time the heart will have become rigid, and whether examined whilst in this state, or subsequently when the rigidity has departed, it is dangerous to assume that the organ is then in the condition which existed at the moment of death, hence the post-mortem condition of the heart is by no means so plainly indicative of the mode of death as is frequently taught.

The subject of post mortem contraction of the heart leads to further consideration of the cadaveric contraction of the skeletal muscles previously mentioned.

Coagulation of the muscle plasma alone is not sufficient to account for all the phenomena which accompany the development of cadaveric rigidity, the shortening undergone by the muscles indicates something more than mere stiffening. Many physiologists are of the opinion that after death rigidity is accompanied by, if not caused by, a true muscular contraction. L. Hermann<sup>1</sup> points out that the chemical processes in vital muscular contraction and in cadaveric rigidity are analogous, with the exception of the formation of myosin from myosinogen, in both,  $\text{CO}_2$ , sarcosolactic, and other acids are formed, heat is evolved, and the muscle current is reversed. He holds that cadaveric rigidity is a contraction of the muscles, occasioned by some unknown stimulation, the contraction lasting longer and passing off much more slowly than an ordinary contraction. Bierfreund<sup>2</sup> divided one ischiatic nerve in recently killed animals, and invariably found that rigidity was delayed on the injured side. Hemisection of the spinal cord below the pyramidal decussation was also followed by delayed rigidity of the side on which the cord was divided. The effect of cutting off the communication with the nerve centres was strikingly displayed in a dog in which the left cerebral cortex was stimulated during life, producing right-sided convulsions, the right half of the cervical cord was subsequently divided, and the animal was then killed. The effect of the convulsions would have been to hasten the onset of rigidity on the side on which they occurred, if no subsequent steps had been taken. The result of cutting off the communication with the brain on that side, however, was that, on examining

<sup>1</sup> *Lehrbuch der Physiologie*

<sup>2</sup> *Arch. für die ges. Physiologie*, 1888



the extremities four and a half hours after, the left side was pronouncedly stiff, whilst the right side was almost as movable as at the time of death. two hours later there was still a marked difference between the two sides. The body of a man who died forty eight hours after an attack of apoplexy also showed delayed rigor on the paralysed side. It was formerly asserted (Nysten) that in hemiplegia there was no difference in the cadaveric rigidity of the two sides so long as the nutrition of the affected muscles had not suffered.

We are now in a position to consider that abnormal occurrence of cadaveric rigidity to which the designation **Cadaveric spasm**, or **Instantaneous rigor**, has been given.

When this phenomenon occurs the last act of life is crystallised in death. It does not consist of an abnormally speedy onset of rigidity after a brief interval of relaxation of the muscles, such as has already been described, but in an absolute prolongation of the last vital contraction of the muscle into the rigidity of death. A few examples will indicate the nature of the phenomenon, and may possibly afford a clue to its interpretation. Palmer, the murderer of Cook, held a handkerchief in one hand at the moment of his execution, it was found tightly clenched in the hand when the body was cut down. Seydel<sup>1</sup> narrates the following cases - (1) A man alighted from a railway carriage in order to procure refreshment, whilst obtaining something to eat, he was disturbed by what he took to be the signal for departure, and ran across the line to his train, not noticing that the locomotive was being backed towards the first carriage. He probably saw the danger when it was too late, and in stooping to avoid it his head was caught between the buffers, and he was instantaneously killed; his arm was outstretched at the moment, and grasped in his hand were some provisions he had just obtained, some hours after, the arm was still extended in the air, and the hand was firmly clasped on the food. (2) By the giving way of a bridge, thirteen persons were suddenly precipitated into the water and drowned. Twelve hours after, when the bodies were recovered, in most of them the extremities were so firmly outstretched as to render it difficult to lay them in their coffins. (3) A man was standing on the ice in the act of lighting a cigar, he fell through, and when the body was recovered, it was found in the upright posture with the cigar and match between the fingers. The body of a man fatally poisoned by water-gas (CO) was found standing upright with one hand grasping the rung of a ladder. Wahneau<sup>2</sup> relates a remarkable case in which the dead body of a woman, aged 44, was found in a standing posture leaning against the outer wall of a house, where it formed a right angle with a closet door, the arms were folded in a natural manner across the chest, and the head was slightly bent forward on to the breast. The woman had been a confirmed drunkard and probably died from some seizure of an uræmic nature, as the kidneys were small and granular. Martin<sup>3</sup> states that a man whom he saw shot dead, fell on the back with the arms stretched forwards and the legs extended in the air, the body immediately became so stiff that it could be moved about by the arms like a piece of wood, the bullet was found to have passed through the base of the skull. Again, Regnard and Loge,<sup>4</sup> giving an account of some experiments made by them on the body of a recently-decapitated criminal, state that two seconds after the head was severed from the trunk the jaws were firmly clenched, four minutes later, the mouth was still firmly closed. Rigidity did not appear in the body until three hours after decapitation, when it began to develop in the lower extremities, six hours after decapitation the arms were still free from any trace of rigidity. The heart—ventricles and auricles

<sup>1</sup> *Vierteiljahrsschr f. ger. Med.*, 1889

<sup>2</sup> *Ibid.*, 1895

<sup>3</sup> *Arch. d'Anthropologie*, 1897

<sup>4</sup> *Le Progrès Médical*, 1887

— continued to beat rhythmically and strongly for twenty-five minutes after decapitation, and the auricles for forty minutes more. One hour after execution the heart was opened, and the left ventricle was found hard and contracted, the right remaining soft.

The coagulation of muscle-plasma, unaccompanied by any other agency, fails to account for this immediate onset of cadaveric rigidity. Some observers have assumed that there is invariably a period of muscular relaxation after death, a line of demarcation between vital action and cadaveric inertia. A glance at the examples just given will show that this assumption is contradicted by facts, the most fugitive muscular relaxation after death would be sufficient to have prevented the occurrence of several of the cases cited. The experiments of Bierfreund tend to prove that some causal relation exists between cadaveric rigidity and the nervous system, that the latter exercises an influence on the onset of the former. Falk and Schroff have demonstrated experimentally that irritation of the medulla produces instantaneous cadaveric rigidity. Seydel groups the cases that have occurred in medico-legal practice into two classes— (a) injuries to the head, with probable irritation of the medulla, (b) toxic action of gases, either CO, or, in those suffocated or drowned, CO<sub>2</sub>. It is suggested that the hypothetical spasm and respiratory centres are in close relation in the medulla and the instantaneous rigidity may result from irritation of these centres. One objection to this hypothesis is that instantaneous rigidity by no means invariably follows death from suffocation by drowning or by hanging, it has been observed that when it does occur from this cause, the individuals have been endowed with powerful muscular development. Whatever the true explanation may be, there is strong reason to believe that the nervous system plays an important part in the causation of instantaneous rigidity. The subject is of great interest to the medical jurist, since the grip of the fingers of a dead body on the handle of a pistol is often the sole evidence in favour of suicide as against homicide.

The **ordinary type** of cadaveric rigidity, as before stated, commences in the muscles of the jaw, and spreads over those of the face, neck, trunk, and limbs. It passes off in the same order—that is to say, the muscles which **first** become **rigid** are the **first** to **lose** their rigidity. Rigidity disappears about the period at which putrefaction commences, and it has been supposed that the disappearance of rigidity is due to the solvent action of ammonia resulting from decomposition of the nitrogenous structures. Myosin is soluble in alkalies, and, according to this view, the ammonia evolved by putrefactive processes dissolves the myosin and thus liberates the muscles. Hermann and Bierfreund, with other physiologists, oppose this view, and state that cadaveric rigidity disappears, or begins to disappear, before any putrefactive changes take place. Hermann attributes its disappearance to solution of the myosin by excess of acid which is formed in the muscles during the continuance of rigidity. In favour of this view, disappearance of rigidity has been observed in muscles retaining a strong acid reaction, it is also urged that an advanced stage of putrefaction would be necessary for the evolution of sufficient ammonia to dissolve the myosin, whereas rigidity often passes off before any obvious degree of decomposition has taken place. Bierfreund examined the muscles microscopically when rigidity had passed off, and found no micro-organisms of putrefaction present, or only isolatedly in the outer layers of the muscles. Halliburton<sup>1</sup> thinks that cadaveric rigidity disappears owing to the action of an unorganised ferment which breaks down the myosin. The inference is that although cadaveric

<sup>1</sup> *Text Book of Chemical Physiology and Pathology*

rigidity passes off about the time putrefaction begins, it is not certain that putrefaction is a necessary factor in causing its disappearance

When cadaveric rigidity has passed away the muscles are soft and flabby they are now dead, and no longer react to electrical stimulation. However short may be the duration of rigidity, when it has disappeared it never returns. There are thus **two periods** of muscular **relaxation** after death, separated by a period of rigidity. During the **first period** of relaxation the **muscles still live**, and demonstrate their vitality by responding to stimulation, during the early part of the period of rigidity they also retain their vitality, but it is latent, or masked, they then lose their vitality, and when rigidity has passed off, and the **second period** of relaxation is arrived at, they are found to be divested of all the characteristics of living matter, in other words, they are in a state of **molecular death**.

### PUTREFACTION.

The last of the phenomena which, under ordinary conditions, follows death, is putrefaction, or the resolution of the organised tissues into their elementary constituents, complex organic bodies, step by step, are resolved into simpler forms, until, ultimately, they are split up into inorganic substances. Putrefaction is the only absolute sign that death has taken place. The agencies concerned in the causation of putrefaction are—*micro-organisms, moisture, air, and warmth*.

**Micro-Organisms.** A large number of organisms are concerned in the process of putrefaction, particularly a group to which Houser gave the name of *Proteus*. These bacteria are ubiquitous, and unless special means are taken either to prevent their access, or to render the tissues unfitted for their use, the dead body invariably succumbs to them. **Moisture** favours the progress of putrefaction. The human body contains a large amount of water in the various fluids and soft tissues, amply sufficient for putrefactive purposes, provided it is not dissipated by rapid evaporation. The addition of moisture from without, especially if accompanied by warmth, greatly accelerates decomposition. The most favourable conditions under which putrefaction takes place are warmth and moisture, with free access of air. The oxygen of the **air** plays its part, but if the air is sterilised it cannot of itself produce putrefaction, as, however, the air invariably contains micro-organisms in abundance, access of air means importation of bacteria. The differing rapidity with which a corpse putrefies in accordance as to whether it is unclad, partially clothed, or completely so, demonstrates the influence exercised by the air in this respect, if a dead body has lain for some time in the open, it will be found that the parts best protected from the air—as the feet are when enclosed in boots—have undergone the least change. **Temperature** is an important factor in determining the rate of putrefaction. A temperature of from 60° to 70° F is favourable to its progress, hence putrefactive changes advance much more rapidly in summer than in winter. When a corpse is interred in the ordinary way, the depth of the grave has an influence on the rate of putrefaction, because the nearer the surface, the greater the effect of the diurnal and seasonal changes of temperature. Except in the case of micro-organisms, excess of any of the conditions which conduce to putrefaction has a contrary effect.—Total submersion of the body in water retards putrefactive changes, excess of air, in the form of a continuous current of air, by

promoting desiccation, tends to dry up the tissues, and thus impedes decomposition, excess of temperature, in either direction, is also unfavourable, a high temperature tends to mummify the soft parts, whilst a very low temperature—below the freezing point—absolutely and indefinitely prevents any putrefactive changes whatever

Thus far the influences considered— which accelerate, and which retard putrefaction— have been of external relation, there remain for consideration those conditions of the body itself which tend to render it more or less prone to decay

**Age.**— The bodies of infants putrefy more rapidly than those of adults. The bodies of spare, old people putrefy slowly. Obesity generally predisposes the body to putrefaction. The mode of death is an important factor. **Diseases** of an adynamic type, as enterica, all septic diseases, and dropsy, death resulting from extensive **mechanical injuries**, from lightning, and from sewer gas, all have a tendency to produce early putrefaction. On the other hand, death from arsenical or from sulphuric acid poisoning may be followed by delayed putrefaction

### SIGNS OF PUTREFACTION.

**External putrefactive appearances** displayed by bodies which have been exposed to the air. The **first outward indication** of putrefaction appears about the **second or third day after death**, in the form of a greenish discoloration on the middle of the abdomen, which spreads to the genital organs, other centres of discoloration of the same character appear on the legs, neck, and back. The eyeball yields to pressure, the cornea is more or less corrugated, and has a milky appearance. At or before this period— four to five days after death— blood-stained fluid oozes from the mouth, together with froth and air bubbles. The gases resulting from decomposition collect in the abdomen, and under the skin, and distend the whole body, the features being unrecognisable. The tongue may be forced between the lips, and the eyeballs protruded. About eight or ten days after death the cornea falls in and appears concave. Still later, bullæ filled with blood-stained serum form on various parts of the surface, the whole body is reddish-brown or greenish, and is still further distended with gas, large areas are devoid of skin, and maggots without number cover the body. The further processes are simply those attending the final dissolution of the soft parts, and are not to be recognised in stages.

Hewitt<sup>1</sup> states that the first changes which occur as the result of bacterial activity are the formation of proteoses and peptone, then leucin, tyrosin, and glycolol, and basic compounds to which the name of ptomine has been given. Next indole, skatole, and phenol, and volatile fatty acids, and lastly, mercaptans, sulphuretted hydrogen, marsh gas, ammonia, carbonic acid, and hydrogen.

Mention may here be made of a special effect, exceptionally produced by the pressure of the gases pent up in the abdominal cavity. In the case of pregnant women dying undelivered it has occasionally occurred that the fœtus has been expelled after the death of the mother. Aveling<sup>2</sup> collected forty four such cases, and from time to time others are recorded. In an example reported by Green<sup>3</sup> a woman aged about twenty died in convulsions (probably uræmic) in the last month of gestation. Fifty three hours after death the dead body of an infant, about term, was found between the thighs of the corpse, and the fundus of the inverted uterus, with the placenta attached, protruded through the vulva, the perineum was considerably torn. No sign of a child was seen when the body

<sup>1</sup> *Manual of Bacteriology*

<sup>2</sup> *Obstetrical Trans*, 1873

<sup>3</sup> *The Lancet*, 1895

of the mother was laid out two hours after death. On an attempt being made to replace the prolapsed uterus it was immediately forced out again by the gaseous pressure within the abdomen, all over the body the putrefactive processes were unusually advanced. In rare cases it appears probable that post mortem parturition may be due to actual contraction of the uterus occurring immediately after somatic death. The musculature of the pregnant uterus, like that of the heart, may retain its molecular vitality sufficiently long to enable this to take place.

The **colour changes** seen in the cadaver are due in the first place to putrefactive destruction of the red blood-corpuscles and to subsequent solution of the hæmoglobin in the serum,  $H_2S$  combines with this forming sulphur-methæmoglobin a solution of which in a thin layer yields a green colour. The altered colouring matter transudes the tissues and tints them accordingly, the anatomical distribution of the veins is indicated by dark lines.

If the body is **buried**, the protection from air afforded by the coffin and the earth in which the coffin is placed has a considerable retarding influence on the rate of putrefaction, the stronger and more air-tight the coffin the longer is the body preserved. This, however, is only markedly the case if the body is placed in the coffin and closed up soon after death, before putrefactive changes have commenced, the retarding influence of sealing up the body in an air-tight receptacle whilst putrefaction is in progress is not nearly so potent as is the case with a fresh corpse. Some kinds of soil preserve bodies which are interred in them, even without coffins, in a remarkable way, dry sandy soil, especially if associated with warmth, promotes desiccation rather than putrefaction, peaty soil retards putrefaction, probably from the presence in it of acids (tannic and others) derived from the vegetable matter of which it is largely composed.

**After death from drowning**, and also in the case of bodies placed in water immediately after death from other causes than drowning, putrefaction takes place more slowly than in air, due to the relatively lower temperature of water as compared with the air above it, and to the exclusion of air from the body. For the first few days little change takes place. Before the end of the first week, the skin on the fingers, the palms of the hands, and the soles of the feet becomes sodden and white, and the face acquires an ashy pallor. The first discoloration takes place on the face, which is swollen and is of a reddish-brown, with patches of green on the eyelids and lips, the discoloration descends to the neck and sternum, sometimes the surface over the latter shows a little redness before the face. The skin is now very wrinkled, in about six or eight weeks the skin of the hands and feet with the nails comes away, often in glove-like form as shown in Plate I, and the hair falls off, or is quite loose. The abdomen is greatly distended with the gases of decomposition, which usually find exit by the natural apertures, the inflation may recur more than once, the body floating with each recurrence, and sinking again when the gases escape. After two to three months, the skin of the arms, thorax, and legs is green. Patches of adipocere will probably be formed on the cheeks and other parts of the body, as shown in Plate II, 1. A month or two later the soft parts are detached, the bones separating subsequently. Harvey Littlejohn<sup>1</sup> directs attention to some striking appearances presented by bodies which are found in the sea. One such appearance is caused by rapid loss of the exposed soft parts of the body, the putrefactive changes in the body as a whole being slower than in fresh water. The rapid disappearance of the exposed parts is chiefly due to the attacks of fish, especially of crabs. This is well seen in Plate II, 2, where the destruction in the face and hands has proceeded to a much greater extent than in the rest of the body. Harvey Littlejohn has seen the whole of the bones of the head, face, hands,

<sup>1</sup> *Edin. Med. Journ.*, 1903

and feet picked as clean as anatomical specimens within ten days of death. Post-mortem injuries are much more frequent, and abrasions of the surface of the body may be bright-red or pink, due to the action of the salt water. In one case, crystals of calcium phosphate, from the size of a millet seed to that of a cherry-stone, were found firmly adherent to the pleura.

It will be observed that the **sequence** of colour changes due to putrefaction **in water** is **different from** that which takes place in **air**. In water the colour-changes begin on the face and spread downwards, in air they begin on the abdomen, and spread downwards and upwards. The order in water obtains whether the body was that of a person drowned (which remains continuously in the water), or whether it was thrown into the water after death, provided that decomposition in air had not previously commenced. If a body is removed from the water in which it has lain for a week or so, it will probably show few signs of putrefaction, a very short subsequent exposure to the air will cause putrefaction to advance at a rapid rate, so that one day in air will be productive of greater change than several weeks further submersion. The rate of putrefaction in water is governed by several conditions, the temperature of the water, the covering of the body, the degree and continuousness of the submersion are all so many factors. Shallow stagnant water is more conducive to putrefaction than deep water, or than shallow water in which there is a current, the influence of the sun-heat is obviously greatest under the first-named conditions. If the body is well protected by clothing, the progress of putrefaction will be delayed. Complete and continuous submersion in deep water is equivalent to low temperature and absence of air, and, therefore, to retarded putrefaction. Dead bodies usually float with the head and extremities below the water-level, the bodies of men in the prone posture, those of women on account of the adipose tissue of the breasts and abdomen—in the recumbent posture in the one case the back, and in the other the abdomen, is at or above the water-level. The bodies of men with much abdominal fat may also float in the recumbent posture. When the body is buoyed up to the surface, the rate of putrefaction is accelerated, especially if the sun's rays are hot, on the other hand, if from any cause, such as the presence of a thick layer of tenacious mud at the bottom of the water, or the entanglement produced by sea-weed, ropes, or other impediments, the body is prevented from floating, the contrary effect is produced.

Caspar gives the following time-ratio for putrefactive changes in *air*, *water*, and in *earth*. Assuming that an equal average temperature obtains in all three cases—one week in **air** equals **two weeks** in **water**, and **eight weeks** burial in **earth** in the usual manner.

**Internal Putrefactive Appearances.**—The internal organs—on account of their inherent differences in density, firmness, proportion of fluid entering into their structure, and in their accessibility to air—undergo putrefaction in a more or less regular order. Caspar's long experience enabled him to formulate the order as follows —

#### **Putrefy rapidly.**

- 1 Larynx and Trachea
- 2 Brain of Infants
- 3 Stomach
- 4 Intestines
- 5 Spleen
- 6 Omentum and Mesentery
- 7 Liver
- 8 Adult brain

#### **Putrefy slowly.**

- 9 Heart
- 10 Lungs
- 11 Kidneys
- 12 Bladder
13. (Esophagus)
- 14 Pancreas
- 15 Diaphragm
- 16 Blood vessels
- 17 Uterus

The mucous membrane of the larynx and trachea is found to be bright or brownish-red or greenish in from three to five days in summer, and in from six to eight days in winter, the only external colour-change at this period will be that of the abdomen. The absence of bony union of the bones of the skull permits easy access of air to an organ feebly resistant to decomposition, hence the earlier decay of the infantile brain in comparison with that of the adult. About the fifth or sixth day after death the stomach exhibits indications of incipient putrefaction in the form of isolated patches of a dirty-red at the fundus, they are first seen in the most dependent part where the post-mortem staining exists. Great caution is necessary in order to avoid mistaking post-mortem changes in the stomach for indications of inflammation, the result of irritant poisons. The spleen may putrefy before the stomach. The liver is usually found firm for several weeks after death. The gall-bladder resists putrefaction much longer. The adult brain shrinks, and the hemispheres soften soon after death, but it takes months, under ordinary conditions, for the brain to melt into the reddish pulp so early seen in the infantile brain. The remaining organs putrefy relatively late. The heart is found fairly fresh when the stomach and liver are in an advanced stage of putrefaction, several months are required to produce in the heart an equal degree of decomposition. The lungs usually show putrefactive changes about the same time as the heart, they may be found in good condition when the external signs of putrefaction are well advanced. The first indication consists of pale red spots of varying size on the surface of the lung, the pleura being raised at these points by the gases of decomposition, these small bullæ are not unfrequently met with even in relatively fresh corpses, but further changes scarcely ever occur until general putrefaction is far advanced, when the colour of the lungs changes to dark bottle-green, and eventually to black, and they subsequently soften and dwindle away. The kidneys resist long, at a considerable interval after death they soften, and become of a blackish-green. The bladder is still more resistant. The œsophagus is much more durable than the rest of the digestive tract. The pancreas does not decay until the body as a whole is much decomposed. The diaphragm may be seen, and its muscular and aponeurotic structures distinguished, four to six months after death. The large arteries last very long, Devergie found the aorta quite recognisable in a body that had been buried fourteen months. The uterus resists the longest of all the soft organs of the body, this enables not only the sex of a cadaver to be ascertained when the external parts are destroyed by putrefaction, but also the occurrence or not of pregnancy or of recent delivery, as was done by Caspar in a foully decomposed body which had lain nine months in a cesspool.

The interval that has elapsed between death and the time that a body undergoing putrefaction is examined cannot be estimated, even approximately, by the stage that the putrefaction has reached, the variations in the degree of putrefaction met with at stated periods after death are so extreme, that the attempt to establish a mean is futile. To take one instance out of many unusually early putrefaction, the following recorded by Taylor and Wilks<sup>1</sup> will serve as an illustration.—A man, aged twenty-six, died in the month of November of typhoid fever and perforation of ileum. Sixteen hours after death there was no cadaveric rigidity, the whole body was bloated, the cellular tissue was so emphysematous that, when the skin was pierced, gas escaped which was easily ignited. The colour of the surface was reddish. The internal organs

<sup>1</sup> *Guy's Hospital Reports*, 1863

were dark, soft, and much-decomposed, and they emitted a very putrid odour. The liver was full of gas.

Abnormally early putrefactive changes do not follow the usual sequence, which, as stated, commences with a green patch on the abdomen. The whole of the surface of the body at once assumes a red colour, with dark lines which show the course of the superficial veins. The appearance is much as though the earlier stages of putrefaction were omitted, or hurried over, so that a condition which usually requires ten or twelve days for its production is arrived at in as many hours.

An illustration of the opposite extreme is related by Taylor and Stevenson<sup>1</sup>. A youth died suddenly, and the body retained such a natural appearance that the friends thought he was in a trance. It was not till thirty-five days after death that they would allow an inspection to be made, when, in spite of the long interval during which the body had been exposed to a warm atmosphere, it was found that putrefaction had made but little progress.

In the face of such extremes, it is clearly impossible to estimate the duration of the interval which has elapsed since death from the stage of putrefaction present in the body.

Some entomological investigations, however, made by Megnin<sup>2</sup> and others, on the **Fauna of the Cadaver**, seem likely to be helpful. It appears that various species of insects, attracted by the successive odours evolved from the decomposing cadaver, severally recognise when the tissues have reached the stage of putrefaction which converts them into a suitable nutrient medium. Megnin found that this sequence of groups of insect life occurs in regular order, and that the lapse of time after death may be estimated by observing the species of insect which is then present. In the first of these groups is the ordinary domestic fly, which lays its eggs on the cadaver in the early days after death. In summer it takes about three weeks for the eggs to develop into the fully formed insect, so that if the chrysalis be found empty, evidence is afforded that more than this period has elapsed since death. The second group comprises flies of the genera *Lucilia* and *Sarcophaga*, which are attracted, three or four days after death, as soon as manifest indications of putrefaction show themselves. A third group consisting of coleoptera (*Dermestes*) and lepidoptera (*Aglossa*) is attracted about three or four months after death by decomposition of the cadaveric fat. A fourth group comprises flies (*Piophilæ* and *Anthomyia*), along with coleoptera (*Necrobia*), which are attracted by the stage of putrefaction arrived at about eight months after death. Through a series of eight groups of insects Megnin traces the progress of putrefaction up to the fourth year after death, when this method of computation comes to an end. Johnston and Villeneuve<sup>3</sup> direct attention to the necessity for observations and experiments on exposed human bodies in each particular locality before entomological data can there be utilised in legal medicine, in their opinion experiments with the bodies of the lower animals are apt to be misleading.

Some of the putrefactive changes in the viscera require to be distinguished from the changes due to inflammation, in the stomach and intestines, for example, appearances may be seen which, though due to putrefaction, might be mistaken for the effects of an irritant poison. The first thing to notice is that the reddening of putrefaction invades the whole of the tissues of which the organ is composed, any variation in this respect is due to the differing density of the structures which enter into the formation of the viscus. Reddening of the stomach due to the action of an irritant during life is limited to the mucous membrane, that due to putrefaction invades the muscular coats as well. After acute gastritis the mucous membrane will easily separate from the structures on which it rests, a condition not met with as a result of putrefaction. The course of post-mortem reddening more clearly follows the distribution of the

<sup>1</sup> *Principles and Practice of Medical Jurisprudence*

<sup>2</sup> *La Faune des Cadavres*, 1894.

<sup>3</sup> *Montreal Med. Journ.*, 1897.



blood-vessels than is the case in inflammatory discolorations. After death the blood-vessels of an inflamed part are filled with blood, the vessels of a patch of putrefactive reddening are empty, or are distended with gas. In serous cavities, fluid may be found which is distinguishable from *intra-vitam* exudation by the absence of inflammatory products—pus corpuscles, shreds of membrane, exudation corpuscles, and also by absence of thickening of the membrane itself.

There are two exceptional processes which may replace ordinary putrefaction **Mummification**, and conversion of the soft parts into **Adipocere**.

### MUMMIFICATION.

This term is applied to desiccation of the soft parts of the body, so that instead of disappearing by colliquative putrefaction, they are dried up and preserved as hard leathery masses, or they may become brittle. In a climate like that of England it is only under exceptional circumstances that mummification takes place. The requirements are dry, warm air, which is most effective when in movement. Mummification under ground is more common in hot countries, in sandy parts. The mummified bodies of newly born infants have been found in boxes or lying exposed to currents of dry air between the rafters of a house and the ceiling of the room below. The length of time required for mummification to take place varies with the conditions of each case, it has occurred in three months.

### ADIPOCERE.

The substance to which this name derived from *adeps, cera-* is applied is an impure ammoniacal soap. The basis is a compound of oleic and stearic acids with ammonia, admixed with which are fibres, and other débris, from organic tissues, with a varying amount of the salts of potash, soda, lime, and iron, in some cases lime replaces ammonia as the base, probably from the presence of much lime in the medium in which the body is placed. The two principal components—fat and ammonia—are derived from the soft parts of the body, the latter resulting from decomposition of the nitrogenous tissues. The bodies of obese subjects and those of children (who usually have much fatty tissue) are more likely to be converted into adipocere than those of spare people. It is probable, however, that the fatty acids concerned in the formation of adipocere may be partly derived from protein as well as from the fatty tissues actually present at the time of death, the appearances found in many advanced cases in which the soft structures of the body have been almost entirely converted into adipocere are inconsistent with the view that the acids are solely derived from the preformed fat. The external conditions which favour the formation of adipocere are submersion of the body in water, and burial in damp soil, or in well-filled graveyards. Cesspools promote either saponification or ordinary putrefaction, sometimes one and sometimes the other. A limited degree of putrefaction occurs before saponification commences.

Adipocere is a waxy-looking substance, having an unctuous feel, it is of less specific gravity than water. Its colour varies from almost white to dark brown, it has a disagreeable odour, especially when heated. If broken, traces of fibres are seen, between the meshes of which the soap is deposited. Cevdalli<sup>1</sup>

<sup>1</sup> *Vierteiljahrschr. d. ger. Med.*, 1906.

points out that this persistence of connective and elastic tissues reveals the anatomical structure of all the saponified organs. Adipocere is a very stable body, and may last for many years.

The time required for the formation of adipocere is naturally variable, depending, as it does, on so many internal and external conditions. It has been held that two or three months' submersion in water is about the shortest time in which it is formed. Traces, however, have been found in from four to five weeks. In moist earth it takes longer to form—eight to twelve months. It may be accepted that in temperate climates indications of saponification in bodies placed under circumstances favourable to its production, may be met with in one month after death, for the whole of the soft parts to be converted into adipocere many years are required.

In hot climates the process may be remarkably rapid. The following cases reported by Mackenzie<sup>1</sup> occurred in Calcutta. (1) A male Hindoo was killed by the kick of a horse and was buried the following day. Four days after burial the body was exhumed in order that an inquest might be held—it was in an advanced state of saponification externally, the heart and liver being also saponified. (2) A young Chinese woman, alleged to have died in childbirth, was buried, circumstances necessitated an inquest, and the body was exhumed seventy-six hours after interment, when it was found to be considerably saponified. These bodies were buried in a soft, porous soil saturated with moisture, the temperature being high, the body last mentioned was enclosed in a wooden coffin. (3) A young European was drowned in the river Hooghly, his body being recovered seven days after. It was in an advanced state of saponification externally, the lungs, heart, liver, kidneys, stomach, and intestines were also saponified, and what is very curious is that the stomach contained undigested food—flesh and potatoes—of which the flesh was entirely saponified, the potatoes not being altered in the least. Other instances of early conversion into adipocere are recorded as occurring in India, in one case, the body was saponified externally and internally in two days.

### CREMATION.

The practice of disposing of dead bodies by cremation is slowly growing in this country. As there is no means of further investigating the cause of death after cremation, the law requires that special precautions shall be taken before cremation, in order to establish beyond doubt that death was due to natural causes.

The regulations governing cremation are laid down by the Home Office under the Cremation Act of 1902. When it is desired that a body shall be cremated, the executor or nearest relative of the deceased must fill up a form of application, which must be returned to the authority undertaking the cremation. Among other things, the applicant is required to state his relationship to the deceased, whether the deceased left any written directions as to the mode of disposal of his or her remains, whether any near relative of the deceased has expressed any objection to the proposed cremation, if so, on what grounds, what was the date and hour of the death, whether he knows or has any reason to suspect that the death was due directly or indirectly to (a) violence, (b) poison, (c) privation or neglect, and whether he knows of any reason for supposing that an examination of the remains of the deceased is desirable. He must also give the name and address of the ordinary medical attendant of the deceased, and the medical practitioners who attended the deceased during his or her last illness. This declaration must be made before a Justice of the Peace or a Commissioner of Oaths. Two medical certificates are required, one from the medical man

<sup>1</sup> *The Indian Med. Gazette*, 1889.

who attended the deceased in the last illness. In this he must state that he attended the deceased before death, and has seen and identified the body after death, whether he is a relative of the deceased, whether, as far as he is aware, he has any pecuniary interest in the death of the deceased, how soon after death he saw the body and what examination of it he made, what was the cause of death, if possible distinguishing between the primary and secondary causes as in the death certificate, whether there was any other cause which contributed to or accelerated death, what was the mode of death, whether syncope, coma, exhaustion, convulsions, etc., whether the deceased underwent any operation during the final illness or within a year before death, if so, what was its nature and who performed it, by whom was the deceased nursed during his or her illness, who were the persons, if any, present at the moment of death, whether, in view of his knowledge of the deceased's habits and constitution he feels any doubt whatever as to the character of the disease or the cause of death, whether he knows or has any reason to suspect that the death or the disease was due directly or indirectly to (a) violence, (b) poison, (c) privation or neglect, whether he has any reason whatever to suppose that a further examination of the body is desirable, and whether he has given the certificate required for the registration of death. The second or confirmatory medical certificate may only be given by a registered medical practitioner of not less than five years' standing, who, in addition, must be either a Medical Officer of Health, Police Surgeon, Certifying Surgeon under the Factory and Workshops Act 1901, Medical Referee under the Workmen's Compensation Act, Physician or Surgeon in a Public General Hospital containing not less than fifty beds, or appointed for the purpose by the Cremating Authority. In his certificate he must state whether he has carefully examined the body, whether he or anyone else made a post mortem examination, whether he has seen and questioned the medical practitioner who gave the first certificate or any other medical practitioner who attended the deceased, or any person who nursed the deceased during his last illness or who was present at the death, and has seen and questioned any relative of the deceased. He must state that he is satisfied as to the cause of death, and must certify that he knows of no circumstances which can give rise to any suspicion that the death was due wholly or in part to any other cause than disease or accident, and that there is no circumstance of any sort known to him which makes it undesirable that the body should be cremated.

The application and certificates are sent to the Medical Referee of the Cremating Authority, who scrutinises them, and, if satisfied that all the requirements of the Cremation Act and of the regulations have been complied with, authorises the remains to be cremated, or he may require a post mortem examination to be made and if this fails to reveal the cause of death, he must decline to authorise the cremation until an inquest has been held and the coroner has given the requisite certificate.

Where it is desired that the body of a person upon whom an inquest has been held shall be cremated, a special form is filled up by the coroner. In this the coroner quotes the verdict to the jury, and states that he is satisfied from the evidence that no circumstance exists which could render necessary any further examination of the remains or any analysis of any part of the body. The coroner's certificate and the application form are then forwarded to the Cremating Authority, the two medical certificates described above not being required.

## CHAPTER VIII

## IDENTITY OF THE LIVING.

THE question of identity may arise both with regard to a living person and also in respect to a dead body. Certain indications serve as clues in both cases, but it is convenient to consider the identity of the dead separately (see Chap. X).

The identification of a living person may come before the Criminal Courts of Law in respect to accusation of murder, assault, and rape, and in the case of escaped prisoners, or of lunatics who have evaded restraint. In the Civil Courts the identity of the claimant to an estate or of a man who has been long absent from the country, and who, on his return, seeks to resume his family relations, may become the subject of legal investigation. Many of the points which can be utilised in civil cases—as family resemblance in features, voice, manner of walking, and other similar characteristics—do not come within the cognisance of the medical expert. His aid is chiefly required to determine the presence or absence of birth-marks, of marks that have been subsequently produced, of deformities, congenital or acquired, of artificial change in the colour of the hair, and other similar matters. He may also be called on to give an opinion as to the possibility of scars or marks on the skin disappearing in the course of time, or by artificial means, without leaving traces, and also as to the alterations in general appearance that may be produced by time, exposure to climatic influence, hardships, and the like. In criminal cases the expert's attention is usually directed to recent wounds, scratches, footprints, and other indications of a struggle, in addition to marks or signs of a longer standing, and also to the presence of hairs, or to stains of blood or other colouring-matter, on the person or clothing. When required to examine the person of an individual who is accused of the commission of an act of criminal violence the practitioner must obtain the consent of the accused and must also explain to him that any indications which may be discovered, of an incriminating nature, will be used as evidence against him at the trial. If the accused will not consent, the examination must not be made. The principle involved is the same as that which governs the verbal interrogation of an accused person. The Law requires that, previous to interrogation, the accused shall be informed that anything he may say in reply may be deposed to in court, and may be used as evidence against him. No one is obliged to furnish evidence against himself, neither by word of mouth nor by submitting against his will to an examination of the person.

**Cicatrices.** The scars left by the healing of wounds differ in size, shape, and character, in accordance with the nature of the injury which gave rise to them. If there has been no loss of tissue, as in the case of a linear incision, and the wound was small and it healed by primary union, the resulting cicatrix may after a time be almost, if not quite, invisible. If the entire thickness of the skin has been cut through, it is doubtful whether the scar ever absolutely disappears, the scars left by venesection and by cupping are, as a rule, permanent and quite evident to the eye.

If a portion of skin throughout its entire thickness is destroyed it is never renewed, the cicatricial tissue which replaces it is permanent, and is wanting in all the characteristics of true skin. It is furnished neither with sebaceous nor sweat glands, nor with hair follicles, and it is sparingly supplied with blood-vessels. In the early stage a cicatrix is of a red colour, changing to brown, subsequently the brown fades, and in course of time the cicatrix becomes white and more or less glistening, which is the characteristic and permanent appearance of old cicatrices. There is no time-limit within which these changes take place, a cicatrix may remain coloured for years. All that can be said is that a red cicatrix is probably not an old one, and that a white cicatrix is not recent, the intermediate stage is of uncertain duration. The visibility of an old linear cicatrix is to some extent dependent on the colour of the surrounding skin, if the skin is white the cicatrix will be little, if at all, different in tint from its surroundings, and may, therefore, escape cursory observation. Advantage may be taken of its relatively feeble blood-supply to demonstrate the existence of the scar, on stimulating the vascular activity of the part by friction, or by slapping it with the hand or with a wet cloth, the normal skin becomes reddened, and the scar, retaining its blanched appearance now stands out in marked contrast. When searching for a small linear cicatrix allow the light to fall obliquely on the part, and examine with a lens. The attempt to obliterate a cicatrix will obviously be unsuccessful. As the skin of the part is absent, and is replaced by another tissue, the removal of that tissue will not restore the skin. A puckered cicatrix may sometimes be dissected out and the edges of the wound brought together, thus forming a linear cicatrix, which materially reduces the amount of disfigurement, the size and shape of the cicatrix will be altered but it will not be obliterated. It is to be noted that cicatrices made in infancy increase in size with the growth of the body, if produced later, they sometimes seem to diminish in distinctness with the lapse of years.

**Birth-marks.** The removal of birth-marks such as moles and *nævi* is usually possible, but not without leaving some trace behind except in the case of very small and superficial *nævi*. If the entire thickness of the skin is not implicated, skilful treatment may cause a small *nævus* to disappear without leaving a mark. It is rare, however, that careful examination with oblique light and a lens fails to yield some trace, although in a general way all indication of the *nævus* may be said to have disappeared. If in the process of removal whether by electrolysis or by the injection of some fluid, one or more of the papillæ under the rete is injured a corresponding depression will be produced on the surface, which, though small enough to escape the unaided eye, will probably be visible if examined, as described, with a lens. Larger or deeper-seated *nævi*, or moles, may be removed by excision, but for the reasons already given, a scar will remain, although different in shape from the original mark.

**Tattoo-marks.** It is a favourite habit of sailors and soldiers, and others, to have various devices tattooed on the skin of the chest, arms, or other parts of the body. The procedure consists in picking out the device, or a portion of it, with needles, and then rubbing in some colouring agent, the operation being repeated until the design is complete. A good deal of swelling and inflammation is produced, and on its subsidence the design is found to be more or less permanently depicted.

The **durability of tattoo-marks** depends on two factors the character of the colouring-matter and the depths to which it is carried.

The **colouring agents** used are carbon in various conditions as gunpowder, soot, Indian ink or lamp black and vermilion, indigo-blue, writing ink, and

other substances. Carbon is permanent as a colouring agent, it is insoluble in all fluids, and its colour does not fade, the other substances named are less likely to be permanent, although they may be so. I have seen a tattoo-mark which was produced forty years ago with ordinary writing ink, and which is still visible. The tattooing in which carbon is used is not black, but is blue-black.

The **position** in which the colouring-matter is located is equally important as regards permanency. If it is superficial, there is the possibility of its removal in time by wear and tear or by artificial means. The reported cases in which tattoo-marks have been erased without leaving any trace were most probably of this class, as shown by the mode in which the obliteration was effected. The application of acetic acid, dilute hydrochloric acid, cantharides, and the like, would remove the epiderm and dissolve out the colouring-matter, or, if it was insoluble, would cause it to be washed out with the exudation from the cutis. Another mode of obliteration of superficial tattoo-marks is to pick out the coloured particles with a needle. In the case of superficial tattooing, it might be possible to obliterate the mark in some such way without leaving any or but very little trace of its former existence. If on the other hand, a permanent colouring-matter is carried down to, or into, the papillæ, it will remain undisturbed for an indefinite period, and cannot be removed without leaving indelible scars. Any attempt to obliterate such a mark by means of acetic acid or by blistering fluid would be futile, and removal with a needle (if practicable) would injure the papillæ, and so lead to depressions on the surface of the skin, which would betray what had been done. Of course, a tattoo-mark, of however permanent a nature, may be removed by excision, or by the application of caustic substances, or of a red-hot iron, but a depressed cicatrix of a size larger than the tattoo will result, in the Tichborne case (1873) some such attempt at obliterating a tattoo-mark above the left wrist had been made. It may be accepted that when tattoo-marks disappear in the lapse of time, or when they can be artificially removed without leaving a trace, either the tattooer, or the colouring agent, or both, are in fault.

**The colour of the hair** may be changed to a darker or to a lighter shade. Hair dyes are usually composed of a solution of a salt of silver, lead, or bismuth, when a speedy result is desired, sulphuretted hydrogen is subsequently used to produce a black sulphide. The metals may be tested for by digesting the hair in nitric acid with the aid of heat, driving off the acid, and then adding the appropriate reagent to a solution of the nitrate in water. Hair may be bleached by means of chlorine, but those who habitually assume light-coloured hair usually resort to an aqueous solution of peroxide of hydrogen. Any mode of bleaching the hair causes it to become lustreless and brittle. The indications by which an opinion may be formed with regard to the naturalness or otherwise of the colour of the hair are that artificially coloured hair has an unnatural hue whether dark or light, and the colour is not uniform, as may be seen by turning the hair over, on careful scrutiny of the hair towards its roots, there will generally be found a difference in tint, especially if the dye has not been applied for a few days. A comparison between the colour of the hair on the head and that on the rest of the body may help in deciding, but too much confidence must not be placed on this, as there is often, naturally, a considerable difference. If no decision can be arrived at for the moment, a day or two passed without the opportunity of applying a dye will place it out of the power of a suspected person to keep up the imposition. Sudden complete blanching of the hair from grief or dread is reported to have occurred, but the evidence is

scarcely conclusive, the hair that grows subsequently, however, may be devoid of pigment Metchnikoff,<sup>1</sup> who attributes the turning of the hair grey or white to the action of phagocytes which swallow up the granular pigment and transfer it elsewhere, believes that increased activity of the hair phagocytes may cause the hair to go grey in a single night

When one or more hairs are found on the person of the accused which resemble the hair of the victim, or *vice versa* hair on the victim resembling that of the accused a careful comparative examination has to be made The points to observe are the tint, diameter, length, and any peculiarities such as are caused by the use of pomade or dye Hair from the head of a woman is slightly thinner than that from the head of a man The hairs from different parts of the body vary in thickness, those from the axilla and from the pubes of both sexes are respectively stouter than from other parts of the body, except from the beard Hairs that have never been cut, as those of the eyelids and eyebrows, taper to a point, hairs that have been cut are either stumpy at the ends or are split into several branches The student is strongly recommended to examine microscopically hairs from the commoner domestic animals the cat, the dog, the horse, the cow, and the sheep, so as to become familiar with their appearance The same advice applies to fibres from various fabrics cotton, wool, silk, linen, etc

**Footprints** sometimes afford evidence of identity The impression may be that produced by the naked foot, or by the boot or shoe In the case of the bare foot, the evidence afforded is rarely conclusive unless there is some distinct peculiarity or deformity Nevertheless, if the impression is sufficiently well marked, a cast should be taken, or in the case of a foot-stain on a hard surface - as when an individual treads with the naked foot into a pool of blood, and then produces a print on the neighbouring unstained part of the floor - a tracing should be obtained by laying a piece of tracing-paper over the dried stain, and carefully going over the outline of it with pen and ink, if possible, the board or flag with the print entire should be removed, as affording still more direct evidence More reliable is the evidence obtained from the impression produced by the tread of a foot wearing a boot or shoe Even in this case it is only when the identical boot or shoe is forthcoming that convincing proof is afforded Conclusions drawn from comparison between the impression and a boot belonging to the accused other than the one that produced the impression are open to serious objection If the impression is sufficiently distinct as to yield a cast, one of two methods may be adopted The impression may be carefully smeared with a flexible feather dipped in oil, and then filled in with plaster of Paris mixed with water to the consistency of cream, ample time must be allowed for the plaster to set before attempting to remove the cast The other method is to hold a large piece of red-hot iron (a cook's salamander is convenient) over the impression, without touching it, until it is well warmed, and then gently to sprinkle the impression with paraffin wax chopped up into a coarse powder, the hot iron should be re-applied after each addition of paraffin, so as to render it fluid, until a sufficiently thick cast is obtained When the paraffin is set, which takes place almost immediately, the cast is carefully removed and any adhering soil gently detached with a soft brush

**Identification by Finger-prints.**—A method of identifying persons by means of the impressions made by their fingers is now widely used in civilised countries The foundations of the present system were laid by Sir William Herschel, who in 1858 required Indian natives to dab their inky fingers on documents in place

<sup>1</sup> *Proc Royal Soc*, 1901

of signatures. In 1890, Sir Francis Galton, in a paper read before the Royal Society, established the fact that the patterns of the impressions are constant throughout life in the same individual. Still later, Sir Edward Henry devised a system of recording the impressions and furnishing a formula for each person which enables any record to be traced without difficulty. The impressions found on the palmar surface of the hand consist of a series of minute ridges and depressions, the ridges being formed by the papillæ of the corium which lie beneath them. The impression is taken by applying the finger, previously washed, to an ink-plate which has been equally covered all over with a thin layer of printers' ink. The finger is rolled on the surface from the radial side of the nail to the ulnar side of the nail. It is then applied to the paper on which the impression is to be recorded, the finger being again rolled in the same way, so as to get a complete impression of the palmar surface of the terminal phalanx and of the transverse creases almost to the end of the digit. At Scotland Yard impressions of the five fingers of each hand are taken.

It is now established beyond all doubt that, though they increase in size, the relative position of the ridges to each other do not alter from birth to death. Further, the pattern in each individual is different, and it has been calculated that the chances are thousands of millions against any two persons exhibiting the same pattern. When the method was first used for criminal purposes, difficulty arose owing to the absence of a satisfactory system of classifying the impressions which would enable any given finger-print to be readily traced in a large collection of other prints. Sir Edward Henry, however, succeeded in classifying the figures made by the ridges into four types—namely, arches, loops, whorls, and composites, and on this basis he was able to devise a formula for each individual which enables the presence or absence of his record in the collection to be determined with the greatest of ease. This system has now very largely displaced the older method of anthropometry of Bertillon, which was based upon various measurements of the human figure.

Dr Garson, who was adviser on identification to the Home Office, has described an ingenious method of identifying invisible finger-prints such as may be left on a weapon or pane of glass by a criminal. A very finely divided powder, such as powdered graphite, is lightly scattered over the surface to be examined, and the latter is then gently shaken to remove superfluous powder. Often, the finger will have left an impression sufficiently greasy to hold fine particles of the powder, and thus a print may be obtained from which identification may be possible. On a dark surface, such as that of a mahogany table, a fine white powder *eq*, calomel may be used. On more than one occasion criminals have been identified by means of such impressions left upon articles they have handled. It is important for the medical jurist to bear this fact in mind, and when dealing with a death from violence to avoid handling more than is absolutely necessary weapons and other objects which might bear impressions, possibly enabling a criminal to be identified.

The finger-print system of identification has become a terror to malefactors, and "high-class" criminals have now taken to wearing gloves while engaged in their activities. Efforts are also made to remove the tell-tale ridges from the fingers, but, short of actual destruction of the skin with production of scarring, this cannot be done. By rubbing the fingers on a rough surface, such as that of a brick, the ridges can be temporarily effaced. If this has been done, however, it is only necessary to remand the suspected person for a few days, when the ridges will grow again.



## CHAPTER IX

**BLOOD AND OTHER STAINS.**

IN medico-legal investigations it is frequently of the greatest importance to determine whether stains on the clothing of the accused, or on a knife or other weapon found in his possession, are or are not due to blood. The question may have to be solved under difficulties, such as result from the stain being very small, very old, or on foul linen. The opinion of the expert only becomes necessary when the stain is of ambiguous appearance, or of microscopic dimensions, the appearance of a fabric which has been drenched with blood is usually sufficiently characteristic as to need little critical examination. For this reason the medical jurist has to select, from the methods used in the physiological laboratory, those which yield the most reliable results with mere traces of blood, the intention is not to demonstrate all the properties of blood, but simply to identify it.

Blood-stains may be examined by three methods **Microscopical, Spectroscopical, and Chemical.**

**MICROSCOPICAL EXAMINATION OF BLOOD-STAINS.**

If a blood-stain is **recent** it will present a bright red or reddish-brown colour, and the fabric on which it exists will be stiffened. The stain should first be carefully examined with a short-focus lens, or with the low power of the microscope. The fabric will be found to have red filaments, and minute clots interspersed in the meshes of its fibres, which present all the outward appearances of coagulated blood. In old blood-stains no coloured substance is seen lying on the fabric as in recent stains. the appearance under the lens is simply that of a fabric that has been stained by some dye. If a small piece of recently stained fabric is cut out and placed in a watch-glass with a few drops of glycerine and water (1 to 10), the colouring-matter will rapidly tinge the solution, and in a short time a few drops squeezed out of the cloth on to a microscope slide may be covered with a thin glass, and examined under a power of 300 to 400 diameters. Red corpuscles will be seen in various conditions in accordance with the age of the stain. If quite recent, many of them will present a fairly normal appearance, others will be contracted and out of shape, irregular in outline, or milled at the edges like coins. With increasing age of the stain the blood-corpuscles shrivel up and become more difficult of recognition, until at last they are completely disintegrated.

Whether the result of the microscopical examination is in favour of the stain being due to blood or not, further investigation is to be made. It is convenient in the first place to have recourse to what is known as the **Guaiacum Test.**

If the suspected stain is on a fabric, a fragment should be cut off and placed on a colour-slab or other white non-absorbent surface. To it add one or two drops of a freshly-prepared solution of guaiacum in alcohol and promote admixture by manipulation with the point of a glass rod, then drop on a little

ozonic ether, or aqueous ether, or aqueous solution of peroxide of hydrogen. If the stain consists of blood, a blue colour is produced, if not, no colour-change takes place. When the stain is on dark-coloured cloth—*e.g.*, from a black coat

the blue colour may be seen on the white surface around the fabric, or a piece of white filtering-paper may be superimposed and pressure applied, in which case the paper will be tinted blue. This test is the most trustworthy when it yields a negative reaction, with the limitation, however, that *very old* blood-stains may not respond to it, although hæmatin readily does so, in its positive phase the production of a blue colour the indication is only to be accepted provisionally, the substance tested *may* be blood, but corroboration is required before a decision is given. Other substances, such as gluten, raw potato, milk, bile, and various oxidizing agents, possess the property of striking a blue colour with guaiacum and peroxide of hydrogen, therefore on no account must a positive opinion be expressed from the indications obtained by this test, even a negative result must not be considered final, as it is an axiom in forensic medicine that every detail demands every possible corroboration. The guaiacum test, therefore, is only a preliminary test which is easy of application, and requires but a fragment of material its use is to pave the way for further inquiry.

**The Benzidine Test.**—Following Schlesinger and Holst, who found benzidine a most delicate means of demonstrating minute traces of blood in faecal matter, M'Queeney has employed the reaction for identifying blood stains on clothing, etc. To half a c.c. of a saturated solution of benzidine in glacial acetic acid, 2 c.c. of hydrogen peroxide are added. This mixture produces a fine blue colour with minute traces of blood, M'Queeney obtaining the reaction in a dilution of 1 in 500,000. The test is open to the drawback that the reagent gives a blue colour with substances other than blood. It should, therefore, only be regarded as a useful preliminary to the spectroscopic test.

The next step is to cut out a portion of the stained fabric close round its margin, and to place it in a watch-glass with a few drops of distilled water. The colouring-matter from a fresh stain will be readily dissolved out, if the stain is old the colouring-matter will have been converted into hæmatin and will be much less soluble. The appearance of the stain, if it is on linen or other light-coloured fabric, will afford an indication of its age. Fresh stains are of a more or less bright red, with the lapse of time the colour becomes browner, but retains a trace of red in its composition for years. The rapidity with which the colour of a blood-stain undergoes this change depends on the freedom with which it is exposed to the air, and upon the presence or absence of chemical substances in the air, such as the oxides of sulphur, hydrochloric acid, ozone, and the like. For this reason blood-stains change more rapidly when exposed to the air in large manufacturing towns than in agricultural districts, hence it is futile to attempt to infer the age of a blood-stain from its appearance, which is altered by external conditions of unknown activity. If the stain appears old, the watch-glass with the portion of fabric covered with water should be protected by an inverted watch-glass, so as to retard evaporation, and left for half an hour or more. It may be preferable to use a test-tube instead of the watch-glass. Some amount of manipulation with a glass rod may be resorted to in order to facilitate solution of the colouring-matter but it is well to avoid mechanical interference as much as possible, lest the resulting solution is rendered turbid. Filtration will remedy this, but if the quantity of coloured fluid is small, the loss from absorption by the filter-paper is of importance. Decantation, after allowing the matter in suspension to subside, is open to the

same objection—loss of fluid, moreover, the suspended matter is often reluctant to subside. Should the stain resist the solvent action of water a saturated solution of borax in cold water may be used, and if very insoluble a few drops of ammonia-water may be added, in which case the solution obtained will be one of hæmatin. Alkalies convert hæmoglobin into hæmatin, but if the blood colouring-matter is insoluble in water it has already undergone this change.

### SPECTROSCOPICAL EXAMINATION OF BLOOD-STAINS.

The fluid thus obtained (after filtration, if necessary) is placed in a small test-tube, or, preferably, in a glass cell with parallel walls, and examined with the **spectroscope**. A small direct-vision spectroscope is convenient, but more exact results are obtained with a single prism table-spectroscope.

Should the amount of fluid for examination be very small, the adaptation of the spectroscope to the microscope, known as the **microspectroscope**, must be employed. In this case it is best to use one of Sorby's cells, which consist of short lengths of barometer tubing about half or three-quarters of an inch—ground to parallel surfaces at the ends, one of which is cemented on to an ordinary microscope slide. A couple of drops of the fluid to be examined will fill the cell, and thus expose a layer of half or three-quarters of an inch in thickness between the source of light and the prisms. It is well completely to fill the cell with the fluid to be examined, and to slide a thin cover-glass on to the upper end of it, avoiding air-bubbles. The cell is then placed on the stage of the microscope, the eye-piece of the instrument being replaced by a direct-vision spectroscope with focussing arrangement. The spectroscopic eye-piece is furnished with a reflecting prism at one side, so that a solution containing blood colouring-matter in a known condition can be made to throw its spectrum alongside that of the blood under examination for the purpose of comparison. When using the microspectroscope, the mirror of the microscope is so adjusted as to project the rays of light through a tubular cell into the instrument, as in ordinary microscopic work—a bright source of artificial light is preferable to daylight, as the presence of Fraunhofer's lines may be embarrassing. The position of the D line can readily be ascertained by bringing into the flame a platinum wire which has been previously dipped in a solution of a sodium salt.

The **spectra** of hæmoglobin and its derivatives are so characteristic that, when obtained, their evidence may be considered conclusive of the presence of blood. For this assertion to hold good, it is essential that the alterations about to be described in the position of the absorption bands should be produced, a final opinion must not be given from an examination of blood colouring-matter in one condition only. There are other substances which yield spectra closely resembling the spectrum of oxyhæmoglobin, but no substance other than blood will give in addition the bands of reduced hæmoglobin and of reduced hæmatin. If all these spectroscopic reactions are produced from a single specimen of colouring-matter, that colouring matter is derived from blood without the possibility of doubt.

The **spectrum of oxyhæmoglobin** (see *frontispiece*) is characterised by the presence of two absorption bands between the D and E lines of the solar spectrum. The first band commences at the D line and extends towards E, the second band commences after a slight gap and terminates at the E line. Some absorption also takes place at both ends of the spectrum, especially at the violet end. It is to be remembered that the appearance of the spectrum, as to breadth of bands and general amount of absorption, varies with the concentration of the solution.

under examination. A good plan is to begin with a strong solution, and gradually to dilute it until the best results are obtained. The spectrum described is the one met with in recent blood-stains. As before stated, the "age" of a blood-stain is not entirely determined by the time that has elapsed since the blood escaped from the body, it may be prematurely aged by the presence of acid vapours in the air, or in a pure atmosphere it may retain its freshness for a considerable time. It is important to bear this in mind.

Very frequently the blood colouring-matter will have passed from the condition of oxyhæmoglobin into that of **methæmoglobin** before the stain comes to be examined. The exact constitution of methæmoglobin has not yet been determined, perhaps the prevailing view is that it consists of hæmoglobin in combination with the same amount of oxygen as that of oxyhæmoglobin, but that the combination is closer, a current of neutral gas— as hydrogen or nitrogen— will not dissociate the oxygen of methæmoglobin as it does that of oxyhæmoglobin. When the stain has passed into the condition of methæmoglobin it will have acquired a brownish hue, and will probably yield an acid reaction. The solution obtained from a stain in this state is less red and more brown than one obtained from a fresh stain, there is however, no great diminution in solubility.

The **spectrum of methæmoglobin** for all practical purposes may be said to resemble that of oxyhæmoglobin with the addition of a thin band in the red, nearer the C line than the D, there is also more absorption of the violet end of the spectrum than is the case with oxyhæmoglobin. The methæmoglobin spectrum then consists of three bands— two in the same position as those of oxyhæmoglobin but paler, and one thin band in the red.

If to a solution of either oxyhæmoglobin, or of methæmoglobin, a reducing agent, such as Stokes' reagent (an aqueous solution of ferrous sulphate with a little tartaric acid, made alkaline with ammonia), or what is preferable ammonium sulphide is added, the spectrum of reduced hæmoglobin is obtained. This spectrum consists of a broad, ill-defined band, occupying very nearly the same position as the two bands of oxyhæmoglobin— that is to say, covering almost all the spectrum between the D and E lines. The absorption of the violet end of the spectrum is about the same as with oxyhæmoglobin, the red end is rather more absorbed.

After identifying hæmoglobin in its two states— in combination with oxygen and deprived of it— some of it is to be converted into hæmatin. Hæmoglobin is decomposed by acids and alkalis into two substances— **hæmatin**— acid or alkaline— which retains the iron of hæmoglobin and **globin**. **Acid hæmatin** has a spectrum not unlike that of methæmoglobin, it is not so easily seen as some of the other spectra of blood, and for its recognition requires a good instrument and a solution of a definite density. **Alkaline hæmatin** also has a spectrum of its own— more difficult to obtain than that of acid hæmatin. It is not necessary for the identification of blood colouring-matter that the two last-mentioned spectra should be examined. It is sufficient if, in addition to the spectra of hæmoglobin or methæmoglobin and of reduced hæmoglobin, the spectrum of **reduced hæmatin**, or **hæmochromogen**, is obtained and identified.

To a little of the original solution of hæmoglobin or methæmoglobin add a few drops of a 20 per cent solution of sodium hydroxide, a change in colour is seen and the original spectrum disappears. To the solution of alkaline hæmatin thus obtained add a few drops of ammonium sulphide, when a further change of colour is at once apparent, the solution becoming somewhat claret-coloured. On examining with the spectroscope the **spectrum of reduced alkaline hæmatin**,

or **hæmochromogen**, the most pronounced of all the blood-spectra, is seen. It consists of two bands slightly nearer the violet end of the spectrum than the bands of oxyhæmoglobin. The first band is dark and is exceedingly well defined at the edges, it is situated about midway between the D and the E lines. The second and broader band, not quite so well defined, commences before the E line extends as far as the *b* line, the violet end of the spectrum is more absorbed than is the case with oxyhæmoglobin. The reducing power of ammonium sulphide is not so active as that of Stokes' fluid, and in consequence, when ammonium sulphide is used, the second band of reduced hæmatin is not, as a rule, fully developed for some time (5 or 10 minutes) after the first, which appears at once. Reduction may be hastened by gently warming the solution after the addition of ammonium sulphide.

If the colouring-matter of the blood-stain has been converted into hæmatin before the fabric is submitted to examination, the first solution obtained from it will yield the spectrum of acid hæmatin. The resemblance between the bands of methæmoglobin and those of acid hæmatin has already been pointed out. The addition of a reducing agent at once distinguishes the one from the other—the methæmoglobin spectrum changes to that of reduced hæmoglobin, the acid hæmatin spectrum to that of reduced hæmatin. When acid hæmatin has been produced by natural causes (without the addition of acid), the use of ammonium sulphide as reducing agent obviates the necessity of previous addition of alkali.

Both hæmoglobin and hæmatin after being reduced can again be oxidised by shaking up the respective solutions with air, to enable this to be done easily, the smallest quantity of the reducing agent must be used that will effect its purpose.

### CHEMICAL EXAMINATION OF BLOOD-STAINS.

If there is sufficient material at command the most conclusive of the chemical tests for blood may be resorted to—the production of **hæmin crystals**. Cut out a piece of stained fabric about the size of a postage-stamp, much less is sufficient with skilful manipulation. Divide this piece into three equal-sized slips and place them one over the other on a microscope slide. Add a minute crystal of sodium chloride, and sufficient glacial acetic acid to saturate the fabric and then a few drops more. Roll a glass rod backwards and forwards over the fabric for a minute or two, so as thoroughly to incorporate the acid with the colouring-matter. A dirty brown fluid can now be pressed out of the fabric by a final passage of the rod, the fabric itself being at the same time withdrawn with a pair of forceps. Bring the fluid to the middle of the slide by stroking it up at each side with the shaft of the rod, and drop on a thin cover-glass. Hold the slide by one end, and pass the centre of it to and fro over the flame of a Bunsen burner, continue until active ebullition, manifested by bubbles rapidly forming under the cover, takes place. Then allow the slide to cool gradually, and when cold examine under a power of 300 diameters.

In the production of hæmin crystals there are one or two precautions to be observed. The amount of sodium chloride must be very small—a crystal the size of a small pin's head is sufficient, if more is added, the field, when examined with the microscope, is seen to be covered with cubes of sodium chloride crystals. With fresh blood, the addition of sodium chloride is unnecessary. Plenty of acid must be used. The boiling demands care—it must be thoroughly done, but if heat is too suddenly applied the glass will probably crack, if excess of

caution is observed— the heat being applied very gradually— the fluid round the circumference of the thin glass will be evaporated, and the dried-up residue will seal in the fluid beneath the cover, so that when ebullition takes place the cover-glass will be projected to a distance

**Hæmin crystals** (*Teichmann's crystals*) are composed of hydrochlorate of hæmatin. They are insoluble in water, alcohol, and dilute acetic and hydrochloric acids, they are soluble in boiling acetic and hydrochloric acids, and in the caustic alkalis. They yield the blue reaction with the guaiacum test, and when incinerated, and the ash is treated with a drop of hydrochloric acid and a solution of potassium sulphocyanide, the presence of iron is demonstrated by the red colour produced. Under the microscope they appear as brown or claret-coloured crystals which have a steely-blue (watch-spring) lustre by reflected light. They usually take the form of rhombic plates, which are frequently superimposed so as to form crosses or stars. The size of the crystals varies— sometimes they are exceptionally large and well-formed, at others they are scarcely recognisable with a power of 300 diameters— when small they occasionally resemble the so-called whetstone crystals of uric acid, but are much smaller. Crystals obtained in the manner described, and possessing the above-mentioned physical and chemical characteristics, afford conclusive proof of the presence of blood.

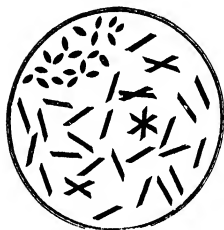


Fig 6—Hæmin crystals

**The distinction of human blood from that of animals** is a question that has long exercised the minds of medical jurists. The blood of birds, fishes, and reptiles presents sufficiently marked differences as to render differentiation between it and human blood possible, the red corpuscles are larger than those in human blood, they are oval in shape, and are nucleated. The only mammals which have blood-corpuscles characteristically different in shape from those of man belong to the camel tribe, which also have oval red corpuscles, but they are not nucleated. The difference in size between the red corpuscles of man

and those of the commoner mammals is so slight that even with fresh blood a witness would not be justified on this ground in deposing positively that a given specimen was derived from a human being, and not from one of the lower animals— much less when the corpuscles have been dried, and subsequently immersed in an artificially prepared substitute for serum. Of the commoner animals the sheep presents the most marked difference in size of red corpuscles as compared with those of man— the proportion is about 4·7 is to 7·7. The following dimensions of the red corpuscles of blood are taken from a table compiled by White<sup>1</sup> from measurements by Treadwell—

Human,	7·940	Horse,	5·503
Dog,	6·918	Cat,	5·463
Rabbit,	6·365	Ox,	5·436
Ass,	6·293	Sheep,	4·745
Pig,	6·101		

Differences in size, which may be of considerable interest from the histological standpoint, are not necessarily sufficiently distinctive for medico-legal purposes. In questions where the life of a possibly innocent man is at stake, all

<sup>1</sup> *The Med. Legal Journal*, New York, 1895

expert evidence must be free from doubt, it may, therefore, be stated that with the exceptions mentioned, it is impossible by microscopical examination positively to distinguish human blood from that of the lower animals. The same attitude is to be taken when the subject of menstrual blood is under discussion, such blood, although possessing certain characteristics as regards coagulability, and reaction to indicators of alkalinity or acidity, is not capable of being differentiated absolutely from blood derived from other parts of the body.

### BIOLOGICAL METHOD OF DISTINGUISHING HUMAN BLOOD FROM THAT OF THE LOWER ANIMALS.

When blood derived from a human being is injected five or six times, at intervals, into the peritoneal cavity of a rabbit, and in a week to ten days after the last injection the animal is killed, the serum yielded by it acts as a precipitant to human blood and (with the exception of monkeys) not to that of other animals. (For methods consult Nuttall<sup>1</sup> Neisser and Sachs<sup>2</sup> and McWeeney<sup>3</sup>)

In 1901, the application of the hæmolsin test to medico-legal practice was first suggested by Deutsch<sup>4</sup> in the same year, Wassermann and Schütz<sup>5</sup> proposed the precipitin test. In experienced hands, the biological test is capable of affording very valuable information, and, in the opinion of many observers, it may be considered to be thoroughly trustworthy, not only with fresh human blood, but also with blood that is undergoing decomposition. Nuttall obtained positive results with human blood which had undergone putrefaction for two months.

Kolmer<sup>6</sup> states that the extract of the blood should be prepared by cutting out a portion of the stained fabric some three inches square. This is torn into shreds with forceps and scissors, not fingers, and placed in 40 c.c. of normal salt solution. If on wood, glass, or metal, the stain should be carefully scraped off. As a control experiment, an unstained portion of the clothing should be extracted in the same way in order to prove that it does not give the reaction. The solution should be stood aside for from two to twenty-four hours, and should not be shaken. The extract should not be stronger than one in a thousand should be colourless by transmitted light, and perfectly clear. These conditions may be obtained by filtering it through a Berkefeld filter. For the preparation of the immune serum, the whole blood may be injected, but it is preferable to use serum only. The serum must be highly potent, sterile, and absolutely clear. The tubes containing the extract and the immune serum should not be shaken. A positive reaction is indicated by a misty cloud at the bottom of the tube in from two to five minutes, which becomes more definite and forms a precipitate in from ten to twenty minutes. Any cloudiness developing after twenty minutes has no significance. The precipitin test is considered by Nuttall, who has had a large experience, to be highly specific. Doubt should only arise between closely allied species—*e.g.*, hare and rabbit, or horse and mule. Kolmer states that "although the blood of the higher apes and even of the lower orders of monkeys may react slightly with human blood, this factor may be determined by observing a proper technique of dilution or the possibility of a given stain

<sup>1</sup> *Blood Immunity and Blood Relationship*, 1904

<sup>2</sup> *Berlin klin Wochenschr.*, 1905

<sup>3</sup> *Lancet*, 1910

<sup>4</sup> *Centralbl. f. Bakteriol.*, 1901

<sup>5</sup> *Berlin klin Wochenschr.*, 1901

<sup>6</sup> *Infection, Immunity, and Specific Therapy*, 1917

being one of monkey blood being definitely worked out" The reaction can be obtained from blood in an advanced state of putrefaction or from a stain that has been dried for a year or more

McWeeney<sup>3</sup> reports two cases in which the test was of assistance In the first a man who was accused of murder was found to have a stain of blood on his cap The precipitin test gave the reaction characteristic of horse blood and not of human blood On subsequent inquiry it was shown that the man was in the employment of a horse dealer and frequently assisted in operations upon horses, especially in the region of the mouth, so that a drop of horse blood might readily have fallen on his cap and dried there The second was also a man accused of murder, and blood was found about his clothes and on his knife He stated that he had used the latter for killing a goat, but on testing, the reaction typical of human blood was obtained While awaiting trial the murderer confessed his crime

**Blood-stains on knives**, or other steel weapons, are frequently very difficult to deal with if the stain is not quite fresh, the colouring-matter has a tendency to combine with the oxide of iron, and thus become very insoluble The saturated borax solution with a few drops of ammonia-water will dissolve out sufficient hæmatin for spectrum analysis, the deposit is scraped off the weapon, and digested with a little of the solution, which is then decanted or filtered from the iron oxide Fresh blood stains on metal are easily dealt with, except when the stain is very thin and in small patches, as after a knife that has been used for homicidal purposes has been wiped with a cloth Careful pencilling of the stained portions with a small camel-hair brush dipped in distilled water or borax solution will generally procure enough colouring-matter for recognition by the microspectroscope If a clasp-knife is undergoing investigation, special attention should be paid to the joint, and if on examination with a lens it appears to contain colouring-matter, the pivots must be withdrawn, and the parts of the knife separated, in this way ample evidence may not unfrequently be obtained

When blood has been projected from a divided small artery, it may reach a neighbouring wall or article of furniture The appearance will probably be that of small splashes or spots, sometimes they take the form of "notes of exclamation" when the jet falls obliquely on the surface Distinction between arterial and venous blood found on clothing, or on furniture, cannot well be made except when a very small artery has sprinkled its contents to a distance in a fine spray Ziemke<sup>1</sup> has made an elaborate investigation into the forms assumed by blood spots according to the direction of the blow, height from which the blood has dropped, and whether or not the injured person was in motion or at rest

### STAINS OTHER THAN BLOOD.

**Rust-stains on knives**, but more especially stains produced by their use in cutting fruit, as oranges or apples—the stain then consisting of citrate or malate of iron respectively—often present a strong resemblance to blood-stains Rust-stains are insoluble in water, and the solutions obtained from the stains produced by vegetable acids on steel yield none of the reactions of blood **Fabrics** may be stained with vegetable, mineral, or anilin colouring-matters Most **fruit-stains** that resemble blood are altered by ammonia-water to a greenish tint, other vegetable-stains are made crimson by the same reagent, some,

<sup>1</sup> International Medical Congress, London, 1913



as logwood, are turned to a bluish-black Dilute acids often alter the colour of vegetable-stains

**Mineral-stains** are often due to the oxide, or some of the salts of iron Ammonium sulphide usually blackens such stains, but as the ammonia of this reagent darkens some vegetable-stains, as archil and logwood, it is necessary to make a control experiment with ammonia-water alone A drop of dilute HCl, followed by a drop of potassium ferrocyanide solution, will yield Prussian blue with iron-stains Most of the **anilin-stains** resembling blood become yellow when treated with dilute  $\text{HNO}_3$ , or tinge the excess of acid yellow In addition to these tests, the absence of hæmoglobin and its derivatives must be proved

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## CHAPTER X

### IDENTITY OF THE DEAD.

THE identification of the dead is required when the body of an unknown person who has died from violence, starvation, or disease, is found in the open air, or in an outhouse, or other uninhabited premises, after a fire in an inhabited house or hotel where strangers are temporarily residing, or after the occurrence of an explosion, or of a railway accident

The difficulties of the investigation are increased if the body is much mutilated, or is far advanced in putrefaction Perhaps the most difficult cases are those in which dismemberment and mutilation have been effected with the special object of preventing recognition of characteristics that might lead to identification

There are certain points to which attention must be directed, which constitute the basis of an investigation having for its object the identification of the body of an unknown person **The stature, age, sex, general state of nutrition, colour of hair, scars, skin-marks** of any kind, **deformities**, indications of past **injury to the bones**, and the presence of **false teeth**, with absence of the natural teeth, all claim attention In the unmutilated fresh body most of these points can be cleared up, the age, as a rule, being the most difficult

When an opinion has to be founded on an investigation of fragments of a body, with the flesh and skin remaining, much depends on the nature of the parts, an entire hand, for example, would convey more information as to the social condition and occupation than a foot If the body has been very much mutilated, and but a few fragments are forthcoming, a part of the thorax or abdomen would probably reveal the sex of the victim Close attention to details is necessary to secure success in all inquiries of this kind When fragments of a body are examined, special attention should be paid to the parts where separation from the rest of the body has been effected An exact description should be written down at the time, and if possible a photograph taken, or a drawing made, so that at a future time, if other fragments are found, it may be ascertained whether they belong to the same body or not

The presence of **scars**, however, small should be noted It may subsequently turn out that the individual whose body is under examination had been bled from the arm, or had received a small wound, the scar of which was known

to have been present up to the time of his disappearance. It has been recommended in cases where there is suspicion of the previous existence of tattoo-marks, to take out the neighbouring lymphatic glands, and to examine them for traces of the colouring-matter.

### THE STATURE.

Within somewhat wide limits, the stature may be determined from the measurement of single limbs, or even of some bones. The length of the arm multiplied by two, with six inches added for each clavicle, and an inch and a half for the width of the sternum, gives the distance between the tips of the middle fingers when the arms are stretched out, and approximately the height of the body (Taylor). The femur equals about  $\frac{2}{3}$  of the body height (Quain). Calculations have been made and tables drawn up with the view of establishing a relation between the length of some of the long bones and the stature of the individual to whom they belonged, but the mean ratio is obtained from measurements ranging over extremes too widely apart to permit of its general application. Estimations of stature from the measurements yielded by one or two bones are very unreliable, if they are given at all it must be in a qualified manner, as they partake largely of the nature of guess-work. When the entire skeleton is available,  $1\frac{1}{2}$  inches are to be added to its length to represent the soft structures.

Criteria for the estimation of **age** have already been given.

If the body is entire, but in an advanced stage of putrefaction, there may be difficulty in ascertaining even the sex. The points to be noted are the hair of the head, that of the pubes, and the development of the breasts. In women the hair of the head is, as a rule, longer and finer than in men, the pubic hair does not usually advance so near the umbilicus in woman as in men. The breasts are fuller in women. In female children all but the first indication will be absent. The clothing will help identification, even if only fragments remain attached to the body. Metallic articles, as rings and other ornaments, may afford a clue. The resistance offered by the uterus to putrefactive changes is to be borne in mind.

If the soft parts have disappeared, the **bones** will afford indications as to **age** and **sex**. When isolated bones, or fragments of bones only, are forthcoming, the first question to be answered is: Are they **human**, or do they belong to one of the **lower animals**? A bone that is intact does not usually present insuperable difficulty, a due acquaintance with human osteology will enable a satisfactory conclusion to be arrived at, for or against. If a bone is not human, it is by no means easy for an ordinary medical man to determine the animal from which it is derived, unless well versed in comparative osteology, it is best for a witness to content himself with the statement that the bone in question is not a human bone. Fragments of the shafts of the long bones may present great difficulties, and demand corresponding cautiousness on the part of the witness. The fragment may be so small and anomalous in appearance that the question becomes—Is it bone or not? The microscope will determine the question by demonstrating the presence or absence of the structural characteristics of bone.

From time to time collections of bones are discovered under circumstances which give rise to suspicion that a murder has been perpetrated. If such a collection is referred to the expert for a report, the first step is to determine whether they are human or not, if they are human, they should be arranged in order, so as to build up a skeleton as completely as possible. It sometimes

happens that duplicate bones are present, which, of course, indicate the dual nature of the remains. A full report should be drawn up, and a drawing or photograph taken when the arrangement is complete. Any abnormality or trace of injury should be noted, taking care not to include under this head the results of violence produced by the implements used in the process of disinterment. The questions to be answered are: What was the **age**, **stature**, and **sex** of the individual or individuals to whom the bones belonged, and are there any indications of the **cause of death**? Each question must be considered separately.

The **age** is to be inferred from the condition of the teeth, and that of the epiphyses of the long bones, together with the other indications previously tabulated. The **stature** can only be arrived at approximately, especially if the skeleton is not complete. The question of **sex**, as regards the skeleton, remains to be considered.

### SEXUAL CHARACTERISTICS OF THE SKELETON.

The **female skeleton** is smaller and of slighter build than that of the male, the individual bones weighing less than the corresponding bones of the male. The following account of the chief distinctions between the bones of the male and female skeleton is taken from Macalister's *Human Anatomy*—

The male skull is larger, heavier, and more ridged than the female, with more prominent mastoid processes, occipital protuberance, zygomatic and superciliary ridges, and a larger

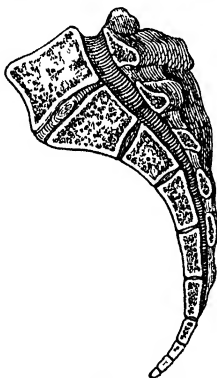


Fig 7 —Male sacrum



Fig 8 —Female sacrum

capacity (11 10), especially in the frontal and occipital regions. The female skull preserves a look of immaturity, and such characters of immaturity as the preponderance of temporal length over frontal, the prominence of the parietal tubera and the concomitant narrowness of the base. The average maximum length from the glabella (the smooth spot between and below the superciliary arches) to the most prominent point of the occiput is 19.6 cm. in the male, and 18.3 cm. in the female. The capacity of the skull—ascertained by filling it through the foramen magnum with No. 8 shot (the other foramina being plugged with cotton-wool) and then measuring the shot in a graduated vessel—averages, in the European male, 1,570 c.c., the European female, 1,378 c.c. The female lower jaw has a less prominent chin, and a weaker, shorter coronoid process than the male, the angle formed by the ramus and body

is greater. The **thorax** is shorter and wider in the female when not distorted by stays. The ribs are more oblique. The twelfth male rib averages 103 mm. in length, that of the female 83·8 mm. The **mesosternum** (body) is more than twice the length of the **presternum** (manubrium) in the male, but less in the female. The **lumbar curve** in the female vertebral column is of greater length, and the lumbo-sacral angle is greater than in the male. The normal adult male sacrum is about 105 mm. high and about 117 broad, the female about 101 high and 118 broad; these relations are expressed by the sacral index  $\frac{\text{breadth} \times 100}{\text{height}}$ . Sacra,

such as those of Europeans, in which this index exceeds 100, are called *platyhiERIC*, those under 100 are *dolichohERIC*, as is the case in most of the black races. The sacral index of British males is about 112·4, of females 116. As a rule, females are *platyhiERIC*.

Ward<sup>1</sup> gives the following characteristics of the curve of the male and female sacrum respectively:—The curvature of the female sacrum occurs chiefly at the lower half of the bone; the upper half is nearly straight. The male sacrum is on an average more curved than that of the female, and its curvature is more equally distributed over its whole length. The male sacrum, in many instances, approximates in form to that of the female, but the female sacrum rarely presents the characters proper to the male. Thus it is much more common to find a straight sacrum in a male subject than one which is very much curved in a female.

The pelvis, as a whole, is the most characteristic part of the skeleton in the two sexes respectively. The male pelvis is deeper, rougher, with deeper iliac fossæ and symphysis, a smaller pubic angle, a vertically ovate obturator foramen, and inflexed tubera ischii.

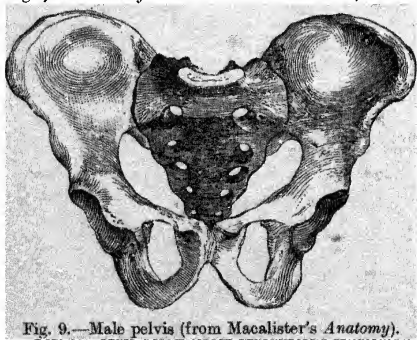


Fig. 9.—Male pelvis (from Macalister's *Anatomy*).

The female pelvis is shallower, wider, smoother, with shallower symphysis, and wider pubic arch, everted ischia, and a triangular obturator foramen. These sex characters are discernible at birth. The sagittal diameter of the brim from sacral promontory to upper edge of symphysis pubis is 100 mm. in the male, 116 in the female. The transverse, 124 in male, 135 in female. The oblique from the sacro-iliac joint to the ilio-pubic eminence, 110 in the male, 126 in the female. The index of the pelvic brim,  $\frac{\text{sagittal} \times 100}{\text{transv. diam.}}$ , varies in different races. The neck of the femur has a lower angle in the adult female than in the male, averaging 118°, the male being 125°.

The skeleton may bear traces of violence inflicted during life. Injuries to the skull caused by firearms are usually too obvious to escape notice. Fractures, especially of the base, might be overlooked except a careful examination is made. Injuries to the cervical vertebræ, from excessive violence due to death by strangulation or by hanging, are to be looked for, also fractures of the ribs. If such injuries have been produced immediately before death there will be no evidence of attempt at repair; the presence of callus shows that an interval elapsed between the receipt of the injury and death. Indications of old fractures should be sought for. The remains of the African traveller, Livingstone, were identified by the presence of an old ununited fracture of the humerus, the existence of which was known to his friends.

<sup>1</sup> *Outlines of Human Osteology*.



## CHAPTER XI

## SUBJECTS INVOLVING SEXUAL RELATIONS.

## SEXUAL ABNORMALITIES

THE subject of sex is one that frequently comes under the notice of the medical jurist. Questions relating to it may arise in regard to newly-born children, as when a woman gives birth to an infant whose sex is doubtful, but which, if a male, is heir to an estate. Later in life sexual development or capacity has to be considered in relation to impotency, rape, impregnation, and allied subjects.

In order that an infant shall be capable of inheriting an estate the law demands that it shall have "the shape of mankind," and if the estate is entailed on male heirs, the question of sex is added. It is impossible to define the limit at which deformity is so excessive as to deprive the subject of it of the right to be considered a human being. The various types of monsters which may be born alive are so diverse that although they may be capable of scientific classification each case has to be separately considered. The law leaves the question open without determining the degree of deviation from the normal which would constitute absence of human shape. Under these circumstances it is useless to attempt any abstract differentiation, all that the expert can do is to make a very careful and detailed examination of the infant, and to report to the court what he has found. It may be observed parenthetically that, contrary to popular opinion, it is criminal to hasten the death of an abnormally formed foetus or monster, although it may obviously be incapable of long survival.

The question of sex in an infant is sometimes equally impossible of determination. Here, however, the morphology of the sexual organs serves as a guide to interpretation. In a living infant, the external generative organs only are available for examination. As an aid to the interpretation of abnormal appearances a glance at the condition of matters at the period of development when the sexes differentiate will be of service. The accompanying two figures diagrammatically represent the male and female external organs in this stage.

In Fig 11 the genital tubercle (*c*) forms the clitoris, the genital folds (*l*) become the labia minora, and the cutaneous folds of the cloacal lips (*L*) remain separate and form the labia majora. The urogenital sinus remains as the vestibule. In Fig 12 the genital folds have approximated and united in the median line, closing on the urethra as far as the glans of the penis, *P*. Those cutaneous folds of the cloaca, which in the female remain separate as the labia majora, coalesce and by their union form the scrotum (*S*). If the genital folds do not unite, the urethra remains open at the raphe (*R*), producing the condition called **hypospadias**.

The conditions which obtain in the fully developed external organs of the female are approximately those which exist before differentiation of the sexes. Further growth and development in a new direction is required to form the male

organs, and during its progress most of the conditions arise which cause ambiguity of sex. It is easy to see that hypospadias may be so extensive as to cause the male organs to appear like abnormally formed female organs. If the testicles have not descended, and the scrotum is cleft, there exists a condition resembling that before differentiation; in such a case it would not be easy to determine the sex from external inspection only. If one or both *testicles* can be found and identified the sex of the infant is thereby determined—it is a *male*. The detection of an *ovary* is equally determinative of the *female* sex. If a glandular organ is accessible it must be carefully examined by the touch, the harder it is the more likely it is to be an ovary. The virgin ovary is largely composed of fibrous tissue, hence it is hard—more so than the testicle. In the adult, compression of the glandular body, if the subject is a male, produces the peculiar sickly, faint feeling which is experienced when the testicles are thus treated. The representative of the penis may or may not be perforate; its prepuce may be free, or it may be continuous with the genital folds or labia minora, the two last-named conditions being in favour of the female sex. If the opening of a cul-de-sac exists below the mouth of the urethra and in front of the bowel, it will probably represent the Mullerian ducts or vagina, and, therefore, is indicative of the female sex; a similar opening, however, may exist in the masculine type of hermaphroditism, being that of the vesicula prostatica—the homologue of the uterus. **Epispadias** consists in the existence of an opening on the dorsal aspect of the penis which is often accompanied by general malformation of the other external genitals; it may be associated with extroversion of the bladder.

After puberty indications may be afforded by the occurrence of functional activity of the sexual apparatus. If periodic discharges of blood take place, they are indicative of the female sex; seminal discharges indicate the male sex. The **bodily conformation** may be taken into consideration—the growth of hair, the general contour, the mammary development together with the vocal intonation. Moral characteristics are not of much value as determinatives, since long custom exercises a powerful influence as regards the sexual proclivities and the habits of the individual.

Developmental malformations of the internal as well as of the external organs of generation are also met with. Individuals so deformed are often, though improperly, called **Hermaphrodites**. The union of the sexes in one being is an old myth having for its basis the assumed power of self-reproduction. In this sense hermaphroditism has no existence; but so far as the linking together in one person of the male and female generative organs, or of parts of them, is concerned, a goodly array of examples is to hand. It is customary to speak of cases of simple malformation of the external genitals which impart a false or ambiguous sexual identity, as cases of *spurious* or *false* hermaphroditism. Such cases have already been described. True hermaphroditism, in which there is an ovary on one side and a testis on the other, or a combined ovary and testis, is exceedingly rare. Blair Bell<sup>1</sup> describes a case in detail, in which the genital glands were removed for supposed carcinoma. One was an ovary and the other an ovo-testis. The patient had a slight moustache, masculine distribution of hair on the body, and a clitoris two inches in length.

<sup>1</sup> *Trans. Roy. Soc. Med.*, 1915.

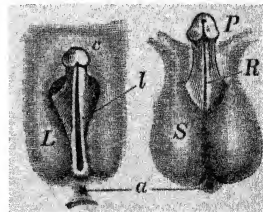


Fig. 11.

Fig. 12.

The following are examples of different kinds of hermaphroditism.—Schmorl<sup>1</sup> describes a case of **lateral and transverse** hermaphroditism in which the external organs had a general resemblance to the male form—there was a well developed but imperforate penis. The scrotum was divided—in the right half there was an organ of the same structure as the testicle, the left half was empty. The internal organs were of the female type—a vagina, which did not open externally, a uterus, with cervix and Fallopian tubes, and an ovary on the left side. Hutchinson<sup>2</sup>, junior, dissected the body of a full grown fœtus, the subject of **transverse** hermaphroditism. The penis was well formed but for a slight hypospadias, the raphe of the scrotum was very well marked. The bulb and prostate extremely well developed, into the latter opened a vagina, above which were perfectly formed uterus, ovaries, and Fallopian tubes. Heppner<sup>3</sup> examined the body of a child two months old, which presented the appearance of **vertical** hermaphroditism. The external organs were of the male type, there was a hypospadiac penis with glans and prepuce, an empty scrotum with raphe, and a prostate gland, but no vasa differentia nor vesicula seminalis. A vagina, a uterus with arbor vitæ in the cervical canal existed, and on both sides round and broad ligaments, well formed Fallopian tubes and ovaries. Under each ovary was a testicle, and near each testicle was an organ resembling the parovarium. Cecccherelli<sup>4</sup> describes a living example of **vertical** hermaphroditism, the subject being fourteen years of age. The mammae were well developed. There was a hypospadiac penis. The scrotum was divided and contained only one testicle, between the folds of the split scrotum, or labia, the neck of the uterus could be felt. The female organs appeared complete. Menstruation had occurred regularly since the twelfth year of age, and the individual had copulated as a woman. From the opening in the penis semen was ejected, a specimen of which was examined by Virchow and found to contain spermatozoa.

**Abnormalities of the Male Organs.**—Individuals in whom one testicle only is in the scrotum, are known as **monorchids**. This condition, whether the result of imperfect development, or of operative interference, is no bar to procreation. Non-descent of both testicles does not necessarily deprive the individual of procreative power, although it is probably obstructive, and it may be a positive barrier, the subjects of this abnormality are called **cryptorchids**. Many post-mortem examinations of individuals labouring under this defect have demonstrated the absence of spermatozoa in the spermatic fluid, but a number of cases in the living have been recorded in which procreative power has existed, so that the positive evidence outweighs the negative. Even absence of both testicles—at least for a time after removal—does not entirely destroy fertility, this, however, can only be due to imperfect ablation, or to the existence of a reserve of spermatic fluid after the organs which secreted it had been removed.

Absence of the **penis**, or of the whole of the external genital organs, may occur either from defective development, injury, or disease. In some cases the penis is united in its whole length to the scrotum, if the organ itself is fully formed, the abnormality is capable of being remedied by surgical operation.

The **female external organs** may be wanting, with or without absence of the **internal organs**. Sometimes the vulva presents a normal appearance, but there is no vagina, which may be either congenitally absent, or it may have been closed by inflammatory processes, not unfrequently from diphtheritic disease in childhood. Surgical procedure can only be resorted to when structures representing the vagina are present. The external organs may be complete, the ovaries and the uterus—one or both—being wanting. The capacity of the vagina may be inadequate for the reception of the penis, usually this condition is capable of being remedied by appropriate treatment. The condition named **vaginismus**, especially if associated with pronounced hysteria, may as effectively bar sexual intercourse as structural atresia of the vagina. Large hernias in either sex may prevent intercourse.

<sup>1</sup> Virchow's *Arch*, 1888

<sup>2</sup> *The Lancet*, 1885

<sup>3</sup> Du Bois Reymond's *Arch*, 1870

<sup>4</sup> *Lo Sperimentale*, 1874



## IMPOTENCE AND STERILITY.

By **impotence** is meant incapacity for sexual intercourse, by **sterility** incapacity for procreation of children. The first term is usually used in reference to the male sex, the second to the female sex, they are, however, respectively referable to both sexes, a man may be sterile as well as a woman, and a woman may be impotent as well as a man. A man may be incapable of procreation, either in consequence of impotence, or, though competent for intercourse, in consequence of absence of the impregnating elements of the semen. A woman may be sterile from absence of the external generative organs rendering her incompetent for sexual intercourse, or, with the presence of these organs, from absence of the ovaries or uterus, in the latter case she would be sterile without being impotent. It will be necessary to discuss the subject with respect to the two sexes separately, and, for convenience, to take impotence and sterility together. In medico-legal practice the subject of impotence and sterility arises in relation to cases of **divorce**, **legitimacy**, and **criminal assaults**.

## SEXUAL DEFECTS IN MALES.

Impotence and sterility in the male may be due to **organic defect** or to **functional disorder**, the organs being anatomically perfect. *Age*, *malformation*, and *constitutional causes* may severally be causal factors.

**Age.**—The two extremes of life may render the individual incompetent for sexual intercourse, no limits can be stated, however, when this occurs.

**Extreme Youth.**—At the age of puberty, which is usually about the fifteenth or sixteenth year, the sexual powers normally begin to develop, capacity for intercourse preceding procreative power. Considerable variation occurs in this respect. The earliest recorded<sup>1</sup> age at which the power of procreation existed in the male is thirteen years, a boy at this age impregnated a young woman. Many cases are recorded of boys becoming fathers at fourteen years of age. Puberty does not consist in a change that takes place at a certain fixed period of life, it is dependent for its early or late occurrence upon hereditary influence, and upon the conditions which surround the individual, morally and physically. For this reason, when examining a case of reputed sexual capacity, the medical man should not be influenced so much by the age of the individual as by his physical and moral development. There are boys of fifteen or sixteen who are sexually mere children, whilst at a much earlier age other boys have the powers and the instincts of adults. The presence of hair on the pubes, and of a developed penis, with or without the general conditions which indicate incipient manhood—as the tone of voice and bodily conformation—are strongly indicative of sexual capacity.

**Advanced age** is no barrier to procreative power. It is true that the sexual capacity diminishes with age, and, consequently, an old man is less likely than a youth to impregnate a woman, this, however, is only stating probability, the medical jurist has to depose to possibility. Active spermatic filaments have been found in the seminal fluid of men of 70 and 80 years of age, and even older, numerous instances have occurred in which men who have married at this time of life have had children. So long as there are active spermatozoa present in the seminal fluid, the possibility of impregnation must be admitted. It may therefore be stated that advanced age is not incompatible with the possible retention of procreative power.

<sup>1</sup> *Brit. Med. Journ.*, 1887

Psychic influences occasionally completely inhibit the sexual emotions. A man from this cause may be utterly unable to have intercourse with one woman although quite competent with another, a perverted sentiment causes the mental attitude to be one of extreme disgust at the idea of coitus with the woman towards whom this sentiment is entertained.

**Debility due to disease** may produce temporary or permanent loss of the sexual functions, most acute diseases of a severe type— as fevers, pneumonia, and the like lead to temporary loss. Of chronic diseases diabetes is one that commonly abolishes sexual energy. Diseases of the **nervous system** have a special tendency to derange the sexual functions. **Tabes** at first increases capacity and later on abolishes it. In **chronic myelitis** the sexual power is usually impaired at an early period, it diminishes more or less rapidly and is finally lost. Ross<sup>1</sup> states that in incomplete paraplegia the sexual power may be preserved for a long time. In the many toxic conditions producing peripheral neuritis, the virile power is more or less impaired, and in some there is complete impotence. **Mumps** sometimes results in metastatic inflammation of one or both testicles, which may lead to atrophy and consequent abolition of procreative power.

**Blows on the head** have been followed by temporary or even permanent loss of the virile power.

The **malformations** of the male sexual organs, described in the previous section, may or may not produce impotence or sterility in accordance with their character and degree. The first consideration is— does the condition prevent the secretion of normal spermatic fluid? and, secondly, if not, is there any absolute mechanical impediment to its being conveyed to the vagina?

**Complete absence** of both testicles renders a man sterile. If they are removed whilst the individual is in full sexual activity, it is possible that one or more fruitful intercourses might take place afterwards, but the power of impregnation is eventually lost. Massazza,<sup>2</sup> experimenting on animals, found that castration does not immediately abolish procreative power, active spermatozoa were discovered in the spermatic ducts nine days after removal of the testicles. The removal of the testicles does not necessarily render a man impotent, although it deprives him of procreative power. History relates that the ladies of imperial Rome were in the habit of taking advantage of this fact by indulging in illicit pleasure with eunuchs, as the act entailed no risk of impregnation. In more recent times the male sopranos (*castrati*), who as a class are barely extinct, have often given rise to scandal. When the testicles exist, although not in their normal situation, the condition imposes no absolute barrier to fertility. Cryptorchids have been known to be fruitful.

**Malformations of the penis** are the chief cause of mechanical impediment to impregnation. **Epispadias** almost invariably makes a man sterile, because the condition renders it next to impossible that the seminal fluid can reach the vagina. **Hypospadias** is or is not a barrier to impregnation in accordance with the degree of malformation and its position, if the penis and scrotum are cleft, or even if the penis only is open at the root, the difficulty is insurmountable by natural means, artificial transference of the semen to the vagina has resulted in impregnation. Since impregnation may result from deposition of semen on the vulva, without propulsion of it into the vagina, great reticence must be observed in expressing an opinion as to the *possibility* of impregnation under adverse conditions on the part of the male. Cases of hypospadias have to be considered on their individual merits, the malformation itself is not a

<sup>1</sup> *Diseases of the Nervous System*, 1883.

<sup>2</sup> *Riforma Medica*, 1891.

fixed condition, and therefore it may or may not render a man sterile. Binding down of the penis to the scrotum is another impediment of varying significance. The same question obtains as with hypospadias—is the deformity of such a character as to forbid totally the possibility of the deposition of semen within the vulva? A curious anomaly may occur with respect to the question of impotence and sterility in the male, the rule is that the former includes the latter, but that the converse is not the case. If a man, by accident or disease, loses the whole of the penis, the testicles remaining, he is obviously impotent, but from what has just been said, he is not necessarily sterile, cases have occurred in which a man thus mutilated has succeeded in effecting impregnation.

### SEXUAL DEFECTS IN FEMALES.

**Impotence and sterility in the female** may also be due to **organic defect**, or to **functional disorder**.

**Age.**—In the female sex puberty usually occurs in this country at fourteen or fifteen years of age. At this age a change takes place in the bodily conformation—the girl becomes more womanly. The commencement of menstruation is the index by which the advent of puberty is recognised, showing that the individual has arrived at the period of life when the capacity of procreation exists. As the development of the virile power in the male may be accelerated or retarded, so in the female may the menses appear early or late. Cases of abnormally early menstruation are not unfrequently recorded, but all discharges of blood from the genital organs of young children are not to be regarded as instances of precocious menstruation. In many cases the discharge is not periodic, and there are no corresponding signs of abnormally early development of the breasts and external genital organs. Exceptionally, however, puberty commences at a very early age, and the reality of the condition is vouched for by the occurrence of pregnancy. The earliest recorded age at which pregnancy occurred in this country is reported by Dodd<sup>1</sup>. A girl began to menstruate at the age of twelve months, became pregnant when eight years and ten months old, and was delivered of a living child which weighed seven pounds. Allen<sup>2</sup> reports a case in which a girl commenced to menstruate at eleven years of age, and at the age of thirteen years and six months gave birth to a fine healthy boy weighing nine pounds, the father of the child was only fourteen years old. In another case reported by Dobson<sup>3</sup> a girl began to menstruate when only eleven years old, she became pregnant at twelve years and nine months, and gave birth to a male child at the age of thirteen and a half. Lefebvre<sup>4</sup> records the case of a girl who was born fully developed with hair on the pubes, she began to menstruate at four years of age, and became pregnant at eight by a man aged thirty-seven, the pregnancy terminating by the expulsion of a mole containing a well-characterised human embryo. Curtis<sup>5</sup> records a case, and after inquiry vouches for the facts, in which a girl became pregnant (by a boy of fifteen) twenty-four days before she was ten years old, she was delivered at the age of ten years eight months and seven days of a healthy child at full term. These cases illustrate the necessity for the medical jurist to determine the presence or absence of puberty by the degree of sexual development attained by the individual rather than by the age. The condition of the breasts, the presence or absence of hair on the pubes, and the general contour

<sup>1</sup> *Lancet*, 1881

<sup>2</sup> *Brit. Med. Journ.*, 1885

<sup>3</sup> *Brit. Med. Journ.*, 1884.

<sup>4</sup> *Gaz. Hebdom.*, 1878

<sup>5</sup> *Boston Med. and Surg. Journ.*, 1863

of the external sexual organs are the indications by which an opinion is to be formed

**Advanced Age.**—Unlike men, women, with few exceptions, are limited in the duration of their procreative capacity, the menopause being usually an indication that the period of fertility is at an end. The average age at which menstruation ceases is about forty-six years, it may cease earlier, or it may be prolonged to fifty, exceptional cases of still later menstruation are recorded as far as and beyond seventy. It is to be observed, however, that as in preternaturally early menstruation, all hæmorrhages from the vagina in advanced life are not to be accepted as evidence of the existence of the monthly function. Uterine polypus and malignant disease of the uterus are conditions which may give rise to hæmorrhage occurring with more or less periodicity. A most extraordinary case of delayed menstruation is recorded by Marx<sup>1</sup>—A woman, forty-eight years old, who had been married twenty years, had never menstruated, at that age she was treated for oophoritis by periodic local abstractions of blood, and subsequently she commenced menstruating at monthly intervals, when the account was published she was fifty-two years old, and continued to menstruate regularly.

Women rarely have children after about forty-five years of age, but exceptions to this rule, and also to the rule that the procreative powers cease at the menopause, are occasionally met with. Davis<sup>2</sup> relates the case of a woman who was delivered at fifty-five years of age, the baptismal certificate of the woman was seen, and her statement of age was thus corroborated. Underhill<sup>3</sup> delivered a woman in her forty-ninth year who had not menstruated for two years. Lavasseur<sup>4</sup> records the case of a woman, aged fifty, who had ceased to menstruate for two years, and who gave birth to a living child at full term. Depasse<sup>5</sup> relates the case of a woman who had ceased menstruating nine years previously, and who had a married daughter forty years old, at the age of fifty-nine years she was delivered of a healthy child, which she suckled, weaning it on her sixtieth birthday.

The form of impotence in men due to **psychical causes** has its complementary condition in women. In men the state is negative—one of pure passivity—in women the impediment is of an active character, and arises from a highly emotional type of hysteria which is associated with vaginismus. A woman in this condition is incompetent for the sexual act, the slightest approach evokes a paroxysm of dread, and of consequent resistance which defies all efforts. Anæsthetics have been used to enable intercourse to be effected, sometimes with permanent success. In other cases the irritability of the vaginal orifice is so intense that the least contact throws the sphincter vaginæ into such powerful contraction as to prevent the introduction even of the finger. In such cases the condition may be more local and physical than general and psychical, and is consequently amenable to surgical treatment.

As intercourse may be accomplished if the woman is entirely passive, it follows that **general disease**—paralysis and the like—does not produce impotence in the female as it does in the male sex. Extreme **bodily deformity** is not necessarily a barrier to intercourse, as women with limbs ankylosed in positions that apparently would forbid all approaches have borne children.

The **local abnormalities** previously mentioned may or may not prevent intercourse. Absence of external genitals, excessive narrowing or absence of

<sup>1</sup> *Przegląd Lekarski*, 1889

<sup>2</sup> *Lond Med Gaz*, 1847

<sup>3</sup> *American Journ of Obstet*, 1879

<sup>4</sup> *Gaz Hebdomadaire*, 1873

<sup>5</sup> *Gaz. de Gynecol*, 1891

the vagina, the presence of a very firm hymen with small aperture, or quite imperforate, adhesion of the labia, tumours filling the vagina, large incarcerated hernias, and other analogous conditions will impede or entirely prevent coitus in accordance with their character and degree. **Absence of uterus and ovaries,** the external organs being complete, will not prevent intercourse, though it will obviously prevent conception. A not unfrequent cause of sterility is the so-called chronic uterine catarrh, in this condition the mucous membrane of the uterus is spongy, and secretes an abnormal amount of clear mucus, which may be blood-stained. Some vaginal discharges by their marked acid reaction render impregnation difficult, spermatozoa are rapidly killed in fluids having a very acid reaction, and also in those which have a very alkaline reaction.

## CHAPTER XII

### RAPE AND UNNATURAL OFFENCES.

**Rape** may be defined as carnal knowledge of an individual by one of the opposite sex without consent, in practice it is understood to refer to carnal knowledge by a male of a female of adult age against her will, or of a girl under a certain age with or without her consent. As the law at present stands —

“Any person who unlawfully and carnally knows any girl under the age of thirteen years shall be guilty of felony. Any person who attempts to have unlawful carnal knowledge of any girl under the age of thirteen years shall be guilty of a misdemeanour. Any person who unlawfully and carnally knows or attempts to have unlawful carnal knowledge of any girl being of or above the age of thirteen years and under the age of sixteen years, or unlawfully or carnally knows, or attempts to have unlawful carnal knowledge of any female idiot, or imbecile woman or girl, under circumstances which do not amount to rape, but which prove that the offender knew at the time of the commission of the offence that the woman or girl was an idiot or imbecile, shall be guilty of a misdemeanour. Provided that it shall be sufficient defence to any charge under sub-section one of this section” (referring to the age of the girl being above thirteen and under sixteen) “if it shall be made to appear to the Court or jury before whom the charge shall be brought that the person so charged had reasonable cause to believe that the girl was of or above the age of sixteen years. Provided also that no prosecution shall be commenced for an offence under sub-section one of this section” (referring to the ages between thirteen and sixteen) “more than three months after the commission of the offence.”—Criminal Law Amendment Act, 1885 (48 and 49 Vict., ch 69)

It is to be noted that the question of age, parenthetically explained, does not refer to idiots nor imbeciles, also that between the ages of thirteen and sixteen an accusation of rape must be made within three months after the alleged offence was committed. Although not stated, it is probably that this time-limit is intended to apply to all ages above thirteen.

A definition of the term “**carnal knowledge**” is necessary for a full comprehension of the above quotation from the Act. By carnal knowledge the law means any degree of sexual intercourse from mere introduction of the penis within the vulva, with or without emission, to complete penetration with emission. If this is done to an adult woman without her consent and against her will, or to a child under any circumstances, it constitutes rape. Formerly

both penetration into the vagina and emission were required, and subsequently (24 and 25 Vict, ch 100) penetration only, with or without emission, was deemed sufficient to constitute the crime. It has been decided by the Courts that introduction of the penis within the vulva, without causing any injury to the hymen, constitutes rape under the conditions above stated. The relation between **age** and **volition** is to be noted. — If a girl of or above the age of **sixteen years** consents to the act, rape is not committed. If a girl of or about the age of **thirteen** and under the age of **sixteen** consents or not, the act on the part of the man is nevertheless criminal, and he is liable to imprisonment for misdemeanour, with or without hard labour, for a term not exceeding two years. If a girl **under** the age of **thirteen** years consents or not, the aggressor is guilty of felony if he effects his purpose, and of a misdemeanour if he attempts only. In the former case the allotted punishment ranges, at the discretion of the Court, from imprisonment for a term not exceeding two years to penal servitude for life, in the latter case the imprisonment must not exceed two years. Therefore, below the age of **sixteen years**, consent or even solicitation on the part of the girl does not do away with the criminality of the act. The same statement applies to women who are **idiots or imbeciles**, consent on their part does not absolve the accused—he is guilty of a misdemeanour. The restrictions of the law apply to prostitutes as well as to women of chaste life, if a man has forcible intercourse with a prostitute against her will, he is guilty of rape, and is liable to be punished accordingly.

If a man, by personating the husband, has intercourse with a married woman with her free consent, he is guilty of rape (48 and 49 Vict, c 69). Formerly this question was left to the judgment of the Court in each case, the consequence being that some judges pronounced the act to be rape, others held that the consent, although obtained by fraud, did away with the criminality of the act. To obviate this want of uniformity the law definitely decides the question as above.

Cases have occurred from time to time that have given rise to the question — Can rape be committed on a woman without her knowledge during sleep? According to the present interpretation of the law, an affirmative answer must be given. The mere introduction of the penis within the vulva would not necessarily cause pain enough to awake a sound sleeper, the act, however, would constitute rape according to the definition of the crime above given. The question was originally propounded when penetration was necessary to establish rape, and at the present time it usually arises in respect to cases in which intromission is alleged to have taken place, when this is the case a qualified answer is required. It would be quite possible for a woman accustomed to sexual intercourse to be violated with complete intromission during sleep. Instances have occurred in which this has taken place, one such is reported in the *Edinburgh Medical Journal*, 1862. Guy<sup>1</sup> mentions the case of a married woman who slept so heavily that her husband frequently had connection with her during sleep. The conditions are quite different, however, when the victim is alleged to have been a virgin up to the time the violence was used. It may be confidently stated that an adult man of average sexual development could not fully penetrate a woman who was a virgin, without her knowledge, during natural sleep. Cases in which it is alleged that this has taken place are *ipso facto* to be treated with grave suspicion, if not with absolute incredulity.

In **abnormal states of sleep**, or stupor, as, for example, *post-epileptic coma*, it is not impossible that a virgin might be ravished without being conscious

<sup>1</sup> *Forensic Medicine*, 1868

of the violence at the time. It is to be expected, however, that on recovery of consciousness she would experience soreness in the private parts, and most probably would find that her underclothing was stained with blood. In Casper-Liman's *Handbuch* a case is narrated of a girl who was subject to epileptic seizures which always left her unconscious for some hours after. A man who was acquainted with this fact on one such occasion took advantage of it, and had intercourse with her without her cognisance. The probability of the occurrence of violation under these conditions would greatly depend on reliable evidence being obtained that the alleged victim was subject to epilepsy followed by stupor.

**Hysteria** and **cataplexy**, as causes of unconsciousness, or of a condition of subjugation to the will of the person accused, are to be received with much caution. It is to be admitted that a girl, or a woman, of a certain neurotic type may for a time be deprived of volition by so-called hypnotism, and whilst in such a condition might be violated without offering effective resistance, or even any resistance whatever. All such accounts, however, are very suspicious, and are—without convincing evidence to the contrary—to be regarded as plausible excuses for non-resistance rather than genuine statements. In giving an opinion respecting such a case, the possibility of the occurrence of hypnotic sleep, or of mental and bodily subjugation induced by suggestion, is to be admitted hypothetically, but great reserve should be maintained in accepting any such condition as accounting for the absence of resistance, or of outcry, whilst the act was being perpetrated.

**Extreme terror** may undoubtedly deprive a girl or woman, for the moment, of the power of offering effective resistance to her violator, volition and physical energy may be temporarily paralysed by the outrageously violent demeanour of an assailant, so that he may accomplish his purpose without any opposition. It is not necessary that the threatened general violence should be directed to the intended victim. In one case recorded by Maschka,<sup>1</sup> the ravisher seized the infant of the woman he desired to violate, and threatened to dash its brains out unless she submitted, the terror thus induced by the appeal to her maternal instincts made the woman submit. In this case, although no physical force was resorted to, the woman was effectively constrained by moral coercion. The law makes no distinction between moral and physical coercion, intercourse against the victim's will, effected after passivity produced by a blow on the head or by threats, is equally criminal. It must be made quite clear, however, that the alleged mental and physical paralysis through fright was actually experienced by the woman, and not feigned as an excuse for offering no opposition. If a woman in the act of illicit intercourse only begins to cry and struggle when she becomes conscious of the approach of a third person, and states that the reason she did not do so before was because she was paralysed by fear, the probability is that she was a consenting party, and that her outcry was prompted by the desire to save her character when detection seemed inevitable. Amos<sup>2</sup> relates a droll case, in which a man was charged with violating a girl, who gave as the excuse for not screaming that she was afraid of waking her mother who slept in the next room.

**Narcotics** or **anæsthetics** are commonly used, and probably much more frequently alleged to have been used, with the object of rendering a woman unconscious and thus placing her at the mercy of her would-be ravisher. The substances usually resorted to for this purpose are—*alcoholic beverages, chloroform, and opium*, or some of its preparations. A woman profoundly intoxicated

<sup>1</sup> *Handbuch*, Bd. 3.

<sup>2</sup> *Lond. Med. Gaz.*, 1831.

would be quite incapable of effective resistance, even if she retained sufficient perception as to be aware of what was taking place. In forming an opinion respecting a case of this kind, the difficulty is to determine whether the woman was as much under the influence of an intoxicant as she professes to have been. A woman who voluntarily drinks an alcoholic beverage until discretion is vanquished by desire, and then yields to the embraces of her male companion, on return to her ordinary state of mind may regret her fall to such an extent as to attempt to visit him with the entire culpability of the act, and so exonerate herself—at least in the eyes of the world—from all fault. In such a case the points for consideration are—The previous character of the woman, her age (a young woman would be more likely to adopt this course than one of middle age), the kind and amount of intoxicant, together with accidental evidence, such as a too clear and detailed account of the transaction being given by the prosecutrix to be consistent with a progressively increasing loss of consciousness. For the reason last named, the time that the man and woman spent together, if ascertainable, might be suspiciously short for an intoxicant of the kind partaken of to produce profound unconsciousness.

With regard to the administration of **chloroform vapour** for the purpose of rendering a woman incapable of resistance, it is to be observed that there are probably only two ways in which this could be done. The first is when a woman *voluntarily inhales* the vapour (which for this purpose includes all anæsthetics capable of being thus administered) for a proper object—as the extraction of a tooth—and when under the influence of the anæsthetic is violated by the administrator. The other way is when the vapour is administered to a woman who is *asleep*. A case of the first kind is quoted at length by Wharton and Stille.<sup>1</sup> A young lady went to a dentist to have a tooth filled, the operation being painful, ether was administered, and, according to the deposition of the prosecutrix, violation was effected whilst she was under its influence. The peculiarity of the case consists in the statement made by the prosecutrix that though unable to resist she was conscious of what occurred to the minutest detail, and moreover could see the disordered state of her clothes when she opened her eyes after the offence had been committed, the dentist having for a moment retired to another part of the room. The young lady discreetly closed her eyes before the dentist returned, and allowed him to readjust her clothing, a few minutes after, she was sufficiently conscious as to discuss the advisability of having the offending tooth extracted on being informed that it was too far decayed to be filled, more ether was given, and the extraction performed. When the effects of the anæsthetic had passed off, she left the house without reproaching the dentist, and did not mention the occurrence until some hours after. To complete the story, the dentist was sentenced to four and a half (!) years' imprisonment, but was subsequently released on representation being made to the authorities that it was quite possible that the whole affair was a hallucination. On reading the detailed account, it is difficult to determine the exact degree of admiration relatively due to the presence of mind displayed by the young lady on the one hand, and the nicely adjusted sentence of the court on the other. This case has been narrated with the object of emphasising a rule that has been many times urged—that a medical man, or a dentist, should never administer an anæsthetic to a female patient without the presence of a third person. This should be one of those rules that has no exception. It is well known that a partial degree of unconsciousness produced by an anæsthetic is not unfrequently attended by delusions, with females such delusions are

<sup>1</sup> *Medical Jurisprudence*, 1860.



occasionally of an erotic type. The vividness of the subjective impression is so great that the person on whom the impression is made thoroughly believes in its objectivity.

The administration of chloroform to sleeping persons in order to prepare them for surgical operations has been tried on many occasions, in some instances the individual passes into the condition of chloroform narcosis without awakening from natural sleep, in others, in spite of very gradual administration, the vapour produces such irritation of the glottis that the subject of the experiment awakes. Dolbeau<sup>1</sup> records some experiments made by him in relation to a case in which it was alleged that a girl had been chloroformed during sleep and afterwards violated. Out of twenty-nine attempts to bring sleeping persons under the influence of chloroform, he succeeded in ten and failed in nineteen.

There is no doubt that a woman once under the influence of an anæsthetic is at the mercy of anyone who chooses to take advantage of her helpless condition. The two ways in which a woman might thus be rendered insensible — by voluntary inhalation and by surprising her whilst asleep — probably comprise all the cases in which violation has been effected with the aid of anæsthetics administered in the form of vapour or gas. Those cases of alleged chloroform narcosis effected by waving a pocket-handkerchief impregnated with an odorous substance before the face, the proceeding being followed by sudden insensibility, are from the nature of things untrue. It is a matter of common experience that the anæsthetist at a surgical operation frequently requires the aid of an assistant during the earlier stages of the administration, to restrain the struggles of a patient who voluntarily submits. Except in the case of a weakly person, it would be no easy matter to administer chloroform single-handed to a woman against her will, the difficulty would probably be quite as great as to commit the rape without the aid of the anæsthetic. A practitioner who is called upon to examine the prosecutrix in such a case should carefully examine the face, neck, shoulders, and wrists for finger-marks or bruises, which would be very likely to be produced (in a genuine case) in overcoming the attempts made by the victim to evade the anæsthetic.

The use of other narcotics to facilitate violation is not common, but the drugging of alcoholic beverages with *opium* or *morphine* has been recorded. The following case of alleged administration of *chloral hydrate* came under the author's cognisance two years after the trial.

A married man, in consequence of the absence of his wife, who was a monthly nurse, was left in his house one night with a servant girl, fourteen years of age. The girl's statement of what happened was that in the evening she complained of toothache, and her master dropped something that tasted like peppermint out of a bottle on to a little cotton wool, and applied it to the tooth, she then went to bed and immediately fell fast asleep. At daybreak she awoke and found her master in bed with her, but he at once got up and left the room. She discovered that she was bleeding from her private parts, which felt painful, she went to sleep again, awoke at the usual time in the morning, got up and did a hard day's work at washing clothes. Although her mistress returned in the morning she made no complaint until evening, when she told her sister the above story. No medical examination was made for five days after the alleged violence had been perpetrated. The following is copied verbatim from the deposition before the magistrates of the medical man who examined the girl:—"There had been penetration, and the nymphæ (*sic*) had been destroyed. There was some inflammation of the private parts. She winced on my touching her. There must have been violence. Chloral would taste like peppermint, and would produce a deep sleep such as would enable any one to commit an act of this sort without the girl knowing."

<sup>1</sup> *Annales d'Hygiene*, 1874

Unfortunately for the accused no rebutting medical evidence was forthcoming. The trial took place at the Manchester Assizes at a time when the public mind was greatly agitated with regard to criminal assaults on young children, an agitation which led to the passing of the Criminal Law Amendment Act of 1885. In spite of the utter improbability of the girl's tale, and of the self contradictory nature of the medical evidence, the prisoner was sentenced to ten years' penal servitude. Two years after, in consequence of representations made to the Home Secretary by the author and by several other medical men, the convict was released. This case admirably illustrates the value and importance of reliable expert medical evidence, and the necessity for every medical witness to know what he is talking about when he is in the witness box. Any well informed medical man would at once have made it clear to the jury that a few drops of a solution of chloral hydrate on cotton wool could not possibly have produced immediate and profound insensibility, lasting for hours, which would enable an adult man to penetrate fully a young girl of fourteen (presumably a virgin) without her knowing. In this way the medical evidence would have served its proper purpose as intended by the law, especially in cases of criminal assault—to corroborate that which is true, and to confute that which is false in the evidence relating to technical subjects given by the prosecution.

By Vict 48 and 49, c 69, the administration of any drug, matter, or thing to a girl or woman with intent to stupefy or overpower, so as thereby to enable any person to have unlawful carnal connection with such woman or girl, constitutes a misdemeanour.

It has been doubted whether a man unaided could fully consummate sexual intercourse with a woman against her will, the woman not being disabled by blows or other violence nor by drugs. The fact that a woman by continued movements of her body could effectively prevent intromission, has led some to deny the possibility of an adult female being violated without aid other than the directly applied physical strength of the man. The question is one which depends on the relative physical strength of the two individuals. A strong man would be likely to succeed with an old or weakly woman, or with a child, a puny man, on the other hand, could be held at bay by a vigorous woman without much difficulty. When the respective muscular developments of the man and woman are more nearly balanced, some discrimination is required in giving an opinion. There is one point with respect to this subject that is often lost sight of—the social position and habits, and the temperament of the woman. Women of the lower classes are accustomed to rough play with individuals both of their own and of the opposite sex, and thus acquire the habit of defending themselves against sportive violence. In the majority of cases such a capacity for defence would enable a desperate woman to frustrate the attempts of her intentioned ravisher. A delicately nurtured woman, on the other hand, is so appalled by the unwonted violence that her faculties may be partially benumbed, and her powers of resistance correspondingly enfeebled.

Under exceptional circumstances the woman's movements may be so hampered as to make her an easy victim. Casper<sup>1</sup> mentions the case of a strong, well-developed woman who was rendered powerless by having her dress thrown over her head, and while thus enveloped was violated by a man single-handed.

It was formerly a matter of debate as to whether rape could be followed by pregnancy. The question needs no discussion, as the impregnation of an ovule is not influenced by volition—it may be accomplished without the participation and against the will of the woman.

Another question has arisen, may injuries to the genital organs produced by rape be sufficient to cause death? An affirmative answer is unfortunately afforded by the occurrence from time to time of cases in which death was directly due to mischief inflicted on the immature sexual organs of young girls by brutal

<sup>1</sup> *Handbuch*, Bd. 1

attempts to effect intromission of the penis, apart from injuries otherwise inflicted Colles<sup>1</sup> reports the case of a girl aged eight years who was violated by an adult She died from peritonitis in six days, and at the autopsy the perineum was found torn and the vagina gangrenous As a result of the early age at which marriage is consummated in India, death of immature girls has not unfrequently occurred from attempts at marital intercourse

### THE PHYSICAL SIGNS OF VIRGINITY.

In the young adult virgin the **breasts** are hemispherical in form, and are firm and elastic to the touch The nipples are small, and are surrounded by areolæ, which are rose-coloured in blondes and darker in brunettes Occasionally a slight secretion of a fluid having some of the characteristics of milk may be found in the virgin breasts this may be due to sexual activity without unchaste habits

The **labia majora** are more or less rounded at their free edges, are firm and elastic, and approximate each other closely the **labia minora** are small and pale in colour The **posterior commissure** is intact Except in rare cases of defective development, the **hymen** in one of its varied forms is always present Its most usual appearance is that of a crescent placed transversely across the entrance of the vagina, with the crescental horns towards the urethra Sometimes the hymen extends all round the orifice of the vagina, and is perforated by a circular or irregularly formed aperture, or by a vertical slit There may be more than one aperture or there may be no aperture at all, constituting the condition known as imperforate hymen The rigidity of the hymen varies as well as its form In some virgins it is patulous and yielding, so that the opening will admit of the introduction of the finger without injury to the hymeneal margin, in others it is firm and unyielding, in which case the passage of any object slightly larger than the opening will tear its free border The **vagina** is narrow, especially in the very young, and its walls are more or less rugose, the rugæ, however, may be replaced by a smooth, non-corrugated surface, due to various conditions which arise from the state of the general health

### PHYSICAL SIGNS OF THE LOSS OF VIRGINITY.

The **breasts** undergo no alteration by a single coitus, and, in the absence of impregnation, little if any by habitual intercourse The **labia majora et minora** only afford corroborative evidence, in the form of an inflammatory condition in recent cases, when much violence has been used The **posterior commissure** is not affected by complete and habitual ordinary intercourse, except in the very young, and in adults when unwonted violence has been resorted to The **hymen** yields the most reliable evidence of loss of virginity, the appearance presented varying with the interval that has elapsed between defloration and the period of examination, it also varies with the age of the individual, the original character of the membrane as to size of aperture and degree of rigidity, the dimensions of the male organ which inflicted the injuries, and the amount of violence brought to bear at the time they were inflicted **Recent injuries** appear as sharp-edged tears or slits which may traverse the entire hymen, and may be continued through the mucous membrane of the vulva and **vagina**, in the ordinary crescentic hymen there is usually one slit in a backward

<sup>1</sup> *Med. Times and Gaz*, 1860

direction towards the commissure, there may be others. In exceptional cases a new aperture is made, usually below the normal opening, leaving a strip of membrane stretching across the mouth of the vagina. From the torn hymen there is usually some hæmorrhage, which is occasionally excessive in any case if the examination is conducted soon after the infliction of the injuries blood-clots or stains will be found on the parts. There will also be an inflammatory condition of the ostium vaginæ and adjacent structures, rendering them painful to the touch, a little later, especially in young children, the inflammatory processes will cause a **mucopurulent discharge** of a yellow or greenish-yellow colour. This discharge is not to be mistaken for the result of gonorrhœal infection which it closely resembles, nor yet for a previously existing leucorrhœa, the distinction is by no means easy, and many false accusations have been made solely on the strength of a discharge which, though due to pathological causes, has been attributed to criminal violence. The discharge which is due to inflammation caused by mechanical irritation, is usually not so copious as that resulting from gonorrhœal infection, from leucorrhœal discharge it is distinguished by indications of an acute inflammatory condition of the mucous membrane which secretes it. Threadworms may cause a discharge from the vagina. Spitzer<sup>1</sup> relates the case of a girl of fourteen who was suspected to have been violated, she had a profuse discharge from the vagina, and on syringing the passage out, threadworms came away, showing the true nature of the case. The presence or the absence of gonococci has been regarded as respectively indicative of the specific or of the innocent character of the discharge. Lober<sup>2</sup> obtained cultivations of gonococci from stains on the clothing in a case of rape. Kratter<sup>3</sup> found gonococci in the vaginal discharge in one out of two cases of rape in which he searched for them, and he attaches much importance to bacteriological investigations in such cases. In medico-legal cases an opinion that gonorrhœa is present should not be expressed unless the organism has been identified both by a microscopical examination and culture methods. The latter alone permits conclusive proof of identity. The rugose condition of the vaginal mucous membrane is not changed by isolated acts of intercourse, and it may be absent in virgins.

If an **interval of four or five days** intervenes between defloration and the period of examination, the appearance of the parts will differ from that above described, especially if the violence has not been excessive. At this period there will be no blood-clots found, and all inflammatory appearance may have disappeared. The sharp edges of the rents in the hymen will be rounded off and the raw surface probably healed. The result is the formation of *caruncula myrtiformes*, which consist of irregularly rounded nodules formed by the remains of the hymen. If recent, the carunculæ are swollen, tender, and of a deep red colour, if more remotely formed, they are firmer, harder, and lighter in colour. It is to be observed that, unless the male organ is disproportionately large, a single intercourse does not result in the formation of a number of nodules representing the pre-existence of correspondingly numerous rents in the hymen. If all the damage sustained by the hymen consists in a single tear, the subsequent appearance will not correspond with what is usually understood by the term *caruncula myrtiformes*, transformation of the hymen into numerous carunculæ is, as a rule, only accomplished by repeated acts of intercourse. Tears in the hymen never unite again, the injury to the membrane is permanent.

<sup>1</sup> *Wien med Wochenschr*, 1892

<sup>2</sup> *Bull med du Nord de la France*, 1887

<sup>3</sup> *Vierteljahrsschr f. ger Med Supplem*, 1891

**The Physical Signs of Virginity or of Non-Virginity in relation to Rape. —**

For the following reasons the signs of loss of virginity do not in themselves constitute proof of rape, nor do the signs of the presence of virginity disprove its occurrence —

**Absence of the physical signs of virginity** may be due to lawful marital intercourse, to illicit but voluntary intercourse, to accidental rupture of the hymen, and, in very rare cases, to congenital absence or imperfect development of the hymen

**The physical signs of virginity may persist** in a female on whom a rape has been perpetrated — from the presence of a patulous hymen with large aperture, especially if the penis of the assailant was of small size, from elasticity of the hymen, which within certain limits may be dilated and may subsequently contract, from extreme youth of the victim, the sexual organs being too small to permit of vaginal penetration (excluding tearing of the parts asunder), and lastly, it is to be borne in mind that mere vulval penetration without the infliction of any injury is sufficient to constitute rape

The hymen may be ruptured by an adequate force of any kind, apart from sexual intercourse. It is reported to have given way from the presence of blood-clots during menstruation, from ulceration following diphtheria or other diseases, from jumping, riding on horseback, or falls on a hard projection. Masturbation has been stated, but probably without sufficient grounds, to be a cause of rupture of the hymen, in the majority of cases of habitual masturbation the hymen will be found intact, the manipulations being limited to the parts anterior to it. Medical examinations or applications may cause injury to the hymen. Some of these reasons for the absence of, or injury to, the hymen are quite feasible, others are far-fetched, each case has to be judged on its own merits.

Complete intromission of the penis may take place without injury to the hymen. The conditions necessary are those previously mentioned — a patulous or elastic hymen, and a not too voluminous penis. Repeated acts of intercourse may take place with no other results than slightly enlarging the aperture through or above the hymen. No inference can be drawn from the presence of a large aperture, the margins of the membrane being intact, as it may be the natural condition of the part. Haberd<sup>1</sup> states that out of forty cases of rape on girls from ten to twenty-eight years of age, in which he examined the genitals of the victims, in fifty per cent. he could detect no anatomical signs of defloration, although the persons implicated admitted that complete intercourse had taken place, and, in some instances, more than once. Distinction must be made between complete intercourse through a patulous hymen, and that form of intercourse which results in impregnation through a small aperture in a tough hymen, which, though a fruitful intercourse, is an imperfect one. Exceptionally, the hymen is both tough and elastic, Maschka mentions the case of a prostitute in whom an uninjured hymen of this kind was present, although complete intercourse had repeatedly taken place.

Rape, including emission, may be perpetrated on **very young children** without injuring the hymen in the least. The external genitals are too small to admit the adult male organ, and the hymen is deeply seated, so that unless sufficient violence is used as to tear open the parts, the assault may be accomplished without leaving more than superficial traces of its commission. A degree of violence may be used on a young child that stops short of lacerating the parts, and is yet sufficiently great to cause subsequent sloughing of the external genitals. Pathological sloughing of the vulva from noma, or from diphtheria,

<sup>1</sup> *Monatsschrift f. Geburtshilfe u. Gynakol.*, 1899

enterica, variola, etc., must be borne in mind in relation to cases of this kind. Anomalous injuries are occasionally produced by sexual violence to young girls. Dorfmeister<sup>1</sup> relates the case of a girl, aged eight years and three-quarters, in whom complete prolapse of the mucous membrane of the urethra was caused by coitus with a boy fourteen and a half years old.

In **married women** there may be absence of the signs of rape, so far as the genital organs are concerned, from conditions the converse of those which obtain in young children. In a woman habituated to sexual intercourse the parts do not sustain injury from a subsequent coitus, even, as a rule, when the act is accomplished with violence, in married women indications of a criminal assault are to be sought for in the presence of spermatozoa in the vagina, and more particularly in manifestations of bruising on various parts of the body.

The circumstances under which the crime is usually committed are such as to render it easy for a designing person to make a charge of rape, and difficult for the accused to rebut the accusation. The crime is one so thoroughly and so universally detested that the victim, or supposed victim, obtains immediate sympathy. It is unfortunately a fact that groundless accusations of rape are very frequent, and in such cases the accused and innocent person suffers from this proneness on the part of the public to accept without question the statements of the prosecutrix. False accusations are not only made by women, and by girls of responsible age, but cases occur from time to time in which mere children are instructed by their mothers to accuse an individual selected for some special reason - extortion of money, or for the sake of revenge - and are not only taught what tale to tell, but are manipulated in such a way as to produce physical indications resembling those caused by criminal assaults, so as to bear out their statements. Fourmier<sup>2</sup> relates the case of a girl, aged eight years, who was said to be the victim of a criminal assault, the person accused being under arrest. Examination of the child showed the presence of violent inflammation of the vulva with erosions of the labia, all the parts being bathed with green pus, the hymen was intact. Several of the glands in the groins were enlarged. After some difficulty it was eventually found out that the mother of the child had produced the injuries by friction with a blacking-brush, an appropriate accusation having been put into the child's mouth to account for them. In cases of this kind it is astonishing with what consistency and pertinacity the child repeats her tale, the threats of severe punishment held out by the mother if she betrays the secret exercise a profound influence in keeping her to the point, if hard pressed the child will burst into tears, and nothing more can be got out of her for the time. In the case just narrated it was only by bribing the child with the promise of a doll that the truth came out, the mother on being confronted with the child's final statement confessed the imposture.

When a rape is committed, as a rule no one is present except the victim and her ravisher, the administrators of the law, therefore, seek for corroborative evidence in addition to the direct evidence of the prosecutrix. The most important corroborative evidence is afforded by medical examination of the prosecutrix, the value of such evidence varies considerably. An early examination of the person of a girl who was intact up to the time the assault was committed yields much more information than one made several days after the commission of the offence on a woman habituated to sexual intercourse. Seeing that vulval penetration (which does not necessarily leave any trace behind) without emission is sufficient to constitute rape, it might be supposed that

<sup>1</sup> *Friedreich's Blätter f. ges. Med.*, 1887.

<sup>2</sup> *Bull. de l'Acad. de Med.*, Paris, 1880.

medical inspection would be futile, in the great majority of cases, however, more than simple contact between the penis and the parts anterior to the hymen takes place, and, consequently, traces of violence are discoverable. Although emission is not necessary to constitute rape, it usually occurs, and traces of the deed are thus left behind. Again, most valuable information is afforded by the discovery of bruises on various parts of the body, caused by the efforts of the ravisher to overcome the resistance of his victim. While due importance is to be attached to the presence of such bruises, it is not always safe to speak dogmatically as to the manner in which they were produced. For example, Reineboth<sup>1</sup> examined a girl aged thirteen, the alleged victim of criminal violence, on whose thighs he found bruises of a peculiar shape which exactly corresponded with the position taken by the tips of the thumb and fingers of each hand if the thighs were grasped as in the act of forcibly separating them, yet the bruises had really been produced by a doubled-up piece of cord with which the girl's father had chastised her whilst she was in bed. Stains of blood or of semen on the clothing are also of importance. Lastly, the presence of gonorrhœa or of syphilis may form a link in the chain of evidence.

### MEDICAL EXAMINATION IN CASES OF SUSPECTED RAPE.

The first duty of the medical practitioner who is required to examine a woman in relation to all matters involving inspection of the sexual organs, is to obtain her free consent. The law holds a woman's person to be inviolable: *an examination made without consent constitutes an indecent assault*. No authority can overrule this privilege, consequently, a medical practitioner who errs with regard to it *cannot shield himself behind an order* received from a magistrate, police authority, or other person. If a magistrate or other authority orders a medical practitioner to examine a woman without her consent, and the order is carried out, both parties are liable to prosecution, a civil action for damages will also lie, as many medical men have discovered to their cost. This question is here emphasised, because medical practitioners are frequently ignorant of the risks they run with regard to the examination of women. It is to be observed that passive submission does not constitute free consent, a woman who simply submits and utters no protest may afterwards deny having consented. The proper way is to make the woman understand the nature and the object of the required examination, and then to ask if she consents to its being made, and on no account to persuade or urge her to submit—her consent must be given of her own free will. When dealing with persons with whom the medical man is unacquainted and in all medico-legal cases, a respectable witness should hear the consent given. If the prosecutrix in a case of rape will not consent to an examination, the medical practitioner must inform the authority from whom he received the order of the fact. In the case of young girls the consent of the father or mother, or of the nearest relative, should, if possible, be obtained before an examination is made.

An examination of the genitals of a female in a case of alleged rape must be very complete. It is to be remembered that the evidence to be given consists not only in a statement of what was obvious, but questions may be asked in relation to what might and ought to have been observed. To enable a thorough examination to be made, the female must be placed in such a position that the thighs may be widely separated, slight injuries to the hymen are rarely sufficiently evident without the membrane being put on the stretch by freely

<sup>1</sup> *Vierteljahrsschr f ger Med*, 1896

separating the labia, the lithotomy position is the best for this purpose. In young children especially, the hymen is deeply situated, and cannot be inspected unless the anterior parts are well stretched open. In recent cases, the parts are so tender that the necessary separation causes much pain, and evokes spastic rigidity of the adjacent muscles, which may seriously embarrass the proceeding, in children this is almost always the case. To wait until the tenderness has subsided is to lose the opportunity of seeing the injured parts in the condition in which they yield the most exact information. A good plan is to apply freely a 20 per cent solution of cocaine hydrochloride five or ten minutes before making the examination, one or two applications, at intervals of three or four minutes, will produce a degree of local anæsthesia which will make an examination possible. If the parts are much lacerated, the cocaine must not be applied too long nor too freely lest toxic effects be produced. *recent raw surfaces are very absorbent*

In cases of recent injury to the hymen the conditions previously described will be found. The presence of carunculæ myrtiliformes in a woman recently violated indicates previous acts of intercourse. All injuries must be carefully noted, and their extent fully determined— as, for example, rupture of the perineum in children, and occasionally in adult virgins to whom excessive violence has been used. The probable amount of blood effused should be estimated from the condition of the underclothing. Specimens of the contents of the vagina should be removed with a glass rod and examined with the microscope for spermatozoa. The surface of the entire body, but especially the thighs, wrists, and neck, should be examined for bruises, scratches, and other marks of violence. Tears, stains produced by mud, paint, or other substances should be sought for on the outer clothing, and, if present, their position noted. The underclothing should be carefully inspected for stains of blood and semen, if the linen is moderately clean, stains produced by semen appear as patches without distinct colour, but of a slightly different tint to the rest of the fabric, which is stiffened at the part stained, a drop of weak gum-water would produce to the naked eye a very similar appearance. After noting their position on the garment all suspicious-looking spots should be carefully cut out and preserved for future investigation, friction or other mechanical disturbance of the stained portion is to be avoided. Amongst the lower orders the underclothing is often exceedingly filthy, being stained with faecal matter, urine, and possibly menstrual blood, in addition to the foulness due to prolonged wear, when this is the case the recognition of seminal stains is difficult. Stains resembling blood-stains should also be cut out for identification. Information should be obtained as to existing or recent menstruation. The presence or absence of venereal disease is to be noted.

The examination of the **dead body** of a female on whom a rape is suspected to have been committed is conducted on similar lines to those followed in the case of the living. The medical witness must be prepared for cross-examination as to the possibility of the appearance indicating bruises being due to post-mortem stains, also as to whether the injuries produced by the rape were sufficient to cause death, whether certain injuries, or wounds, were inflicted before or after death, as well as the means probably used to inflict them. If first examined on the spot where it was found, attention must be paid to the position of the body as a whole, and to that of the limbs and clothing. The surroundings must be investigated for indications of a struggle, especially if the body was found out of doors, foot-marks scrutinised, and, if necessary, casts of them taken in the manner described on p 55. Any peculiarities of the



soil—as sandy, clayey, and the like—are to be noted, and if of a markedly characteristic appearance, a specimen should be preserved for comparison with the soil possibly present on the boots of an accused person. The state of the weather at the time the crime was committed is also to be ascertained as affording a possible clue in case of the arrest of a suspected person. The subsequent examination of the body should be of such completeness as to enable the witness to testify to the presence or absence of fractures of the bones (especially of those of the skull), the condition of the internal organs, and of the natural apertures of the body as regards the presence in them of foreign bodies. When death has been caused by suffocation (a not unfrequent cause of death in fatal criminal assaults), ecchymoses, due to the asphyxia, may be present in various parts of the body. Lediard<sup>1</sup> directs attention to the possibility of such hæmorrhages being found beneath the mucous membranes of the vagina and of their being mistaken for signs of local violence—the presence of vaginal ecchymoses should cause the practitioner to search for corresponding extravasations in other parts of the body.

If the accused is in custody a medical examination will probably be required of his person and clothing. That which has already been said as to the necessity for consent on the part of a woman to an examination of this kind, applies with equal relevance to a man. No one has the legal right to make an examination of the generative organs of a man, nor to order such an examination to be made, against his will. It is further to be observed that if an examination of any part of the body of the accused, whether of the genital organs, or of the limbs, or the trunk, for bruises, scratches, or for the presence of hair, derived from the victim, is deemed advisable, not only should consent be obtained, but, before making the examination, the accused should be informed that any indications discovered, which may tend to incriminate him, will be made use of at the trial.

Except when venereal disease has been communicated to the violated female, and the discovery of a similar disease on the accused would constitute a link in the chain of evidence, examination of the male, as a rule, yields little information. If spermatozoa are found in the urethra, or on the shirt, their presence may be accounted for by the avowal of recent intercourse with another woman than the one violated, or of an emission without intercourse. If, as is most likely, no spermatozoa are found in the urethra, recent emission would not be disproved, since the act of urination would wash all traces away. The points to notice on the male are—the presence of spermatozoa on the person or clothing, of blood-stains (in a case of recent rape on a virgin, traces of blood should be specially sought for under the prepuce and about the frænum), the condition of the outer clothing as to tears and stains of various kinds, the boots with any adherent soil or mud, recent scratches on the face or other parts of the body, the presence of long hairs on the coat or trousers (if found they should be preserved for comparison with the hair of the victim), together with a careful observation of the general build of the man, so as to enable an estimate to be made of his physical strength—whether he appears strong enough to have overcome a woman such as the prosecutrix.

### EXAMINATION OF SEMINAL STAINS.

If a drop of mucus is taken from the vagina in order to ascertain whether spermatozoa are present or not, all that is necessary is to place the mucus on a slide, to cover it with a thin cover-glass and examine with a power of 300 to

<sup>1</sup> *Med.-Chirurg. Trans.*, 1896

400 diameters. If semen has been deposited on a fabric and has dried, the spermatozoa require softening and liberating from the surface to which they are attached by the inspissated albumen of the fluid. If pure water is used the spermatozoa tend to swell out to such an extent as to cause them to disintegrate, the heads separating from the filaments. To justify a statement on oath that a given specimen contains spermatozoa it is essential that one or more should be seen in a complete form, for although the appearance presented by the heads and the filaments apart is very suggestive, it does not constitute positive evidence, it is essential, therefore, that precautions should be taken to minimise the risk of separation of head and filament. The first precaution is to avoid handling the fabric more than necessary, and to keep the stained portions flat, without creasing them, the next is to use a fluid for softening that will not cause the spermatozoa to swell too much. Ungar<sup>1</sup> recommends for this purpose a very dilute solution of hydrochloric acid in distilled water—one drop to 40 cubic centimetres, a few drops of the acidulated water is placed in a watch-glass with a small strip of the stained fabric so that its lower end dips into the fluid. The time requisite for softening varies from a few minutes to several hours in accordance with the age of the stain. It is well thoroughly to soften the albumen before attempting to detach the spermatozoa or else they will be broken. If only the lower end of the strip of cloth dips into the acidulated water, there is little risk of the spermatozoa being detached prematurely, therefore it is better to allow too much rather than too little time, old stains require four to five hours. When the softening is complete, the fragment of fabric is removed with a pair of dressing forceps and gently dabbed on a microscope slide. The deposit thus obtained is covered with a thin glass cover and examined as previously directed. As the spermatozoa are very translucent objects, it has been proposed to stain them with the view of making them more visible, either the dry or the moist process may be adopted. Ungar recommends a double stain for the dry process, which may be thus used—After a strip of fabric has undergone the requisite softening in the acidulated water, it is withdrawn with the forceps, allowed to drain a little, and then dabbed on a thin glass cover so as to leave a deposit, which is allowed to dry at the temperature of the surrounding air. When dry the cover is taken up with the forceps and rapidly passed deposit side upwards—over a Bunsen flame so as to thoroughly harden the deposit. When cold, the cover-glass is floated—deposit side downwards—on a solution of eosine in a watch-glass, the solution is composed of eosine 2.5 c. gm., alcohol 30 c. c., and distilled water 70 c. c. The watch-glass with its contents should be protected so as to retard evaporation, and allowed to remain one hour, after which the cover-glass is removed, drained, exposed until dry, and then lightly washed with one part alcohol to two parts water. The next step is to place the cover-glass in a solution of logwood (Friedlander's) for a period varying from a few minutes to an hour or more, the time depending upon the action of the stain, which is variable. The preparation is then washed and examined in the usual way. The results obtained are—that the back part of the head is stained dark blue, and the front part, with the middle piece and filament, an intense red. A simpler and more practicable method is to combine the softening and staining solutions in one. A solution of methyl green .15 to 3 gm. in 100 c. c. of water, to which from 3 to 6 drops of hydrochloric acid are added, in accordance with the amount of colouring material used, will serve this purpose. The strip of fabric with the seminal stain is allowed to dip into a few drops of this dye in a watch-glass and to remain for several hours.

<sup>1</sup> *Vierteiljahrsschr. f. ger. Med.*, 1886

It is then removed with the forceps and dabbed on a slide, the deposit may be examined at once in the moist state

A human **spermatozoon** consists of a head, to which is attached a short rod-like piece continued by a long slender filament, the whole measuring 0.05 mm in length. The head is egg-shaped when viewed in one direction, and more pointed—like the outline of the flame of a candle—when viewed sideways. The whole spermatozoon is composed of transparent protoplasm with a delicate fibrillous outline. With the amplifying power used for the purpose of identification—300 to 400 diameters—no trace of structure is visible. Spermatozoa retain their activity within the sexual organs of women for many hours, they have been found active in the vagina seven to eight days after emission. They preserve their form after cessation of activity for almost an indefinite time, and may be recognised by the methods above described in a seminal stain several years old.

Florence<sup>1</sup> has devised a chemical test for semen. A drop of a strong aqueous solution of iodine and potassium iodide (the iodine being in excess) is allowed to come in contact with a drop of the fluid obtained by soaking a seminal stain in a little water, a precipitate is at once formed which under the microscope is seen to consist of crystals extremely like haemin crystals both in form and colour. Johnston<sup>2</sup> examined with this test the seminal secretions from a number of cadavers and obtained positive reactions in all but two in which the disease of the seminal vesicles and prostate existed, spermatozoa, however, being present. Posner<sup>3</sup> has obtained the reaction in cases of azoospermia and also with an extract of ovarian tissue. Cardile<sup>4</sup> states that cholin and a number of the alloxan and xanthine series gives Florence's crystals, which may be obtained therefore from urine. Bocarius<sup>5</sup> found that the spermatozoa of the lower animals and other substances of a seminal nature, vegetable as well as animal, yield Florence's crystals. He gives a full bibliography of the subject. Florence's test for semen occupies the same position as the guaiacum test does for blood—it is merely a preliminary test. Berberio's<sup>6</sup> test consists of a saturated solution of picric acid which, when added to human semen, or to a seminal stain, produces crystals of a peculiar appearance like Charcot-Leyden crystals. The semen of the ordinary domestic animals, and the secretions, excretions, and exudates of the human organism, do not give the reaction. Levinson<sup>7</sup> obtained the reaction from a stain between three and four years old, but he found that advanced putrefaction was inhibitive. The reaction is not only given by semen which contains spermatozoa, but also by that in which spermatozoa are absent, though less markedly and with less certainty. It is doubtful to which of the components of semen the reaction is due, probably to the prostatic secretion.

The parts of the female underclothing on which a seminal stain is most likely to be found after violation are the front and back of the chemise in the neighbourhood of the genital organs. It is to be noted that a stain may be due to seminal fluid and yet no spermatozoa may be present, spermatozoa are sometimes absent in the semen of an individual at one time and present at another. Very exceptionally certain trichomonads, known as *trichomonas vaginæ*, have been found in the vaginal mucus, and the fact of their existence has been utilised to weaken evidence given as to the presence of spermatozoa in cases of alleged rape. A glance at the representations of spermatozoa and these trichomonads (Figs 13 and 14) is sufficient to show the difference between the two, which is so great that one could not possibly be mistaken for the other by any competent observer. The head of the trichomonad is circular in outline, it is much larger than the head of a spermatozoon. It is granular in place of being structureless, and the filament or tail is proportionally very much shorter.

<sup>1</sup> *Arch. d'Anthrop. Crim.*, 1896.

<sup>2</sup> *Boston Med. and Surg. Journ.*, 1897.

<sup>3</sup> *Berliner klin. Wochenschr.*, 1897.

<sup>4</sup> *Arch. di Farm. e Terap.*, 1899.

<sup>5</sup> *Viertejahrschr. f. ger. Med.*, 1901.

<sup>6</sup> *Med. Leg. Journ. of N. York*, 1905-6.

<sup>7</sup> *Berliner klin. Wochenschr.*, 1906.

Spermatozoa have such a characteristic appearance that when found intact—that is, with head and filament continuous—the stain from which they are derived may, without hesitation, be pronounced to be due to seminal fluid. It is to be observed, however, that occasionally an object strangely resembling a spermatozoon may be seen in a microscopic specimen of such a nature as to

preclude the possibility of a spermatozoon being present, it is, therefore, advisable in cases of alleged rape to identify more than one zoosperm.

**Blood-stains** on the under-clothing of females alleged to have been violated are to be examined after the methods described on p. 57. When blood-stains are present, the blood may not have entirely permeated the fabric on which it was deposited, should this be the case, note



Fig. 13 --Human Spermatozoa



Fig. 14 -- Trichomonas Vaginæ

whether the stain is on the *inner* or the *outer* surface of the fabric in relation to the body of the wearer. Maschka<sup>1</sup> mentions a case in which he found the stains to have been produced by the blood of a bird, the imposture being subsequently confessed. Romberg<sup>2</sup> mentions an extraordinary blunder made by a medical man, who stated that certain stains on a little girl's chemise were respectively caused by blood and semen, although they really resulted from the child having eaten a plum tart in bed—the supposed blood-stains being due to plum juice, and the seminal stains to fatty matter derived from the pastry.

### UNNATURAL SEXUAL OFFENCES.

Under this head are comprised *Sodomy*, *Pederastia*, *Bestiality*, and *Incest*.

**Sodomy** means unnatural sexual intercourse between two human beings usually of the male sex. The converse form **Tribadism**—gratification of the sexual instinct between two human beings of the female sex—is not an offence against the law in this country. **Pederastia** is that form of sodomy in which the passive rôle is played by a boy, the active agent being man or boy. **Bestiality** means sexual intercourse between mankind and the lower animals. These offences are dealt with in section 61 of 24 and 25 Vict., ch. 100, thus—

“Whosoever shall be convicted of the abominable crime of buggery, committed either with mankind or with any animal, shall be liable, at the discretion of the court, to be kept in penal servitude for life, or for any term not less than ten years.”

Much new light has been thrown upon the etiology of sexual perversions by the work of Freud, Ferenczi, and others who have followed in their footsteps. According to this school, the normal sexual instincts of the adult are only attained after the individual has passed through a series of phases which begin in early childhood. Freud recognises the beginnings of sexuality in the pleasurable gratifications of certain desires, and he even includes in these the pleasure of sucking the mother's breast and pleasurable sensations attending defæcation which are experienced by the infant in early months. Later, when its interest is aroused towards others, the child passes through a bi-sexual

<sup>1</sup> *Handbuch*, Bd. 3

<sup>2</sup> Casper Luman, Bd. 1

stage, and its affections may be directed towards persons of either sex, but most often in the case of boys towards the mother and of girls towards the father. In one phase the love of the child is centred upon itself, and this is very frequently associated with masturbation long before the attainment of puberty. With normal development these phases are passed through and eventually the natural inclination towards the opposite sex develops. In some cases, however, the normal progress is arrested in one of the earlier stages, and when adult life is reached the tendency is still towards the form of gratification desired at that particular stage. Then, one of two things may occur. As a result of social teaching, the individual declines to recognise in himself these tendencies which differentiate him from other people, and does not even admit them to consciousness. The conflict which then ensues between his conscious and subconscious minds is likely to lead to the development of a psychoneurosis. The further consideration of this condition pertains to books on the neuroses. On the other hand, the individual may admit to consciousness his abnormal tendencies, and endeavour to gratify them, with the result that he may bring himself into conflict with the law.

The etiology of homo-sexuality is really more complex than would appear from the simplified statement above, and it is probable that more than one psychic trend results in this condition. Ferenczi<sup>1</sup> has distinguished two kinds of homo-eroticism in males. The "active" homo-sexual feels himself a man in every respect, is, as a rule, energetic, and does not display effeminacy in his bodily or mental characters. It is only that the object of his inclination is changed. The other, or passive type, appears to have the psychic make-up of a woman, is attracted by mature powerful men, and regards women with antipathy or even hatred. He is effeminate in his manner, may let his hair grow long, and is often of a girlish build physically. Ferenczi regards this as a true sex intermediate stage. Other forms of perversions can also be traced to infantile fixations. Exhibitionism is the term applied to a practice often observed in tiny children of displaying with pride their genital organs. In adult life the persistence of this tendency may lead to offences such as indecent exposure of the person.

There is now a large volume of literature dealing with sexual perversions, and further information as to their origin, manifestations, and treatment by psycho-analysis must be sought for in appropriate books. The medical jurist is only interested in these questions when there have been actual breaches of the law. The general principles of the Freudian school have been to a large extent accepted, though there is diversity of opinion in many details, and it is sufficient to say here that the whole trend of opinion in recent years has been towards regarding sexual perverts and inverters more as persons suffering from abnormal psychic development and needing treatment by the physician than as depraved persons whose proper custodian is the gaoler.

It is probable that inverted sexual feeling is much more widely spread than is supposed, the cases that come before the courts of law being but the shadow of the reality. Not many years ago the police of a town in the North of England took possession of a public room in which a ball of a very exclusive character was taking place. It was found that the company consisted wholly of men, half of them being dressed as women, in order that their proceedings should not be observed, they had provided themselves with blind musicians.

The passive as well as the active agent in the commission of sodomy is amenable to the law, except when overcome by force, or by being drugged,

<sup>1</sup> *Contributions to Psycho-analysis*, translated by Ernest Jones, 1916

or when under fourteen years of age. The act is criminal when committed with a woman as well as with a man. As in rape, the passive agent can give evidence as to the perpetration of the crime. The medical evidence in cases of sodomy is not nearly so important as it is in cases of rape. The active agent shows no indication of the habit, although the contrary has been asserted. The passive agent, if long habituated to the act, may display certain indications which are by no means constant. The sphincter ani may be relaxed, and the anal opening more patent than normal, a condition which may be accompanied by prolapse of the mucous membrane, a smooth condition of the skin round the anus is sometimes met with. Lacerations in the sphincter ani, or of the mucous membrane of the rectum, are only found in recent cases, and are more common in young boys than in adults, in such cases there may be swelling and inflammation present. Unless obtained from within the anus, the discovery of spermatozoa is only important if the passive agent is very young, as in older subjects they may emanate from the individual himself. With regard to the distinction between the active and the passive agents, it is to be remembered that the rôles are interchangeable: the agent that on one occasion takes the passive may on the next take the active part, for this reason, when making an examination in a case of alleged sodomy, it is well to investigate the condition of both the parts concerned in each of the agents. A medical man told the author that when acting as assistant to a police surgeon, he was on one occasion required to examine a man and a boy who were in custody on the charge of sodomy. He examined the man solely as the active agent, and the boy as the passive agent, without finding any indications, at the trial it turned out that the boy and not the man was the active agent.

The possibility of sodomy being perpetrated on a man against his will, or during sleep, is contrary to common sense. With regard to young boys, it is possible that they might be overcome by force. The same remarks apply to this crime as to drugging and other modes of taking away the power of resistance, as were made with regard to rape.

The Criminal Law Amendment Act of 1885 enlarges the scope of the law, with respect to the commission of indecent offences by two or more male persons, beyond the previous enactments relating to sodomy, Part I, section 11 (48 and 49 Vict. ch. 69), runs thus —

“ Any male person who, in public or private, commits, or is a party to the commission of, any act of gross indecency with another male person, shall be guilty of a misdemeanour, and being convicted thereof shall be liable at the discretion of the court to be imprisoned for any term not exceeding two years, with or without hard labour ”

Medical evidence in cases of bestiality is limited to a search for hairs (derived from the abused animal) on the clothes and person of the accused.

**Incest.**—The punishment of Incest Act, 1908, makes it a misdemeanour punishable by not less than three nor more than seven years' penal servitude, for a male person to have carnal knowledge of his grand-daughter, daughter, sister, half-sister, or mother. It is immaterial that the carnal knowledge was had with the consent of the female. Further, any female of or above the age of sixteen who with consent permits her grandfather, father, brother, half-brother, or son to have carnal knowledge of her is guilty of a misdemeanour, punishable by not less than three nor more than seven years' penal servitude, or by imprisonment not exceeding two years with or without hard labour. Proceedings under this Act must be held *in camera*.

**Indecent exposure** of the person is a criminal offence which does not come

under the notice of the medical jurist in the way that the above-mentioned offences do, but his opinion may be asked as to the criminal responsibility of the offender. Certain forms of cerebral degeneration dispose those who are subject to them to exhibit their persons in public. In this country the cases have been mostly those of elderly men developing senile dementia, or in younger men, the subjects of epileptic automatism, of alcoholism, or of the initial stage of general paralysis. Krafft-Ebing<sup>1</sup> has directed attention to a form of sexual psychopathy in which there is a special psychical degeneration occurring in the early years of life, mostly referable to hereditary influence or to pathological conditions, such as rachitis. He gives illustrative cases in which there is an apparently uncontrollable tendency on the part of the subjects to expose their persons to others. Among these is one by Freyer<sup>2</sup> of a man, thirty-five years of age, who was arrested for lingering about a girls' school, and when he succeeded in attracting the attention of the pupils exhibiting his person. For the same offence he had been in prison more than half a dozen times before, for periods of from three months to three years, on the last occasion he was very properly sent to a lunatic asylum instead of to prison. He had been subject to epileptic seizures, and it is worthy of remark that when the impulse to expose himself developed, the epileptic attacks ceased.

## CHAPTER XIII

### THE SIGNS OF PREGNANCY AND OF DELIVERY.

MEDICAL practitioners may be called on by legal authorities to ascertain the occurrence or non-occurrence of pregnancy under the following circumstances -

- (a) When a woman sentenced to death pleads pregnancy as a bar to execution,
- (b) when a woman whose husband is recently dead asserts that she is pregnant with an heir to the estate, the heir-at-law, to protect himself from fraud by the importation of a spurious heir, may demand proof of pregnancy. In both these instances the ancient proceeding was to empanel a jury of twelve matrons or discreet women to make the necessary investigation and to report to the court. The law still requires this procedure formally to be observed, but the jury is now assisted by a medical practitioner, who makes the actual examination.
- (c) When a woman who has been seduced claims increased damages on account of being with child, (d) when an unmarried woman, or widow, or a married woman living apart from her husband, is, as she alleges, libellously accused of being pregnant, or (e) when a woman who has lost her husband through culpable neglect of some person or persons, claims damages for his loss and for the future support of an unborn child.

The caution previously given respecting medical examinations of women is to be borne in mind when the practitioner is about to investigate cases of suspected pregnancy, abortion, or natural delivery. Not only must consent be obtained, but the suspected person should be informed that any incriminating appearance revealed by the examination may be used as evidence against her.

<sup>1</sup> *Wiener med. Blätter*, 1892

<sup>2</sup> *Zeitschr. f. Medizinbeamte*, 3 Jahrg.

## SIGNS OF PREGNANCY.

An exhaustive description of all the various indications of the existence of pregnancy which are of value to obstetricians, is out of place in a treatise on Forensic Medicine. What the medical jurist has to depose to on oath is not the probability or otherwise of a certain woman being pregnant, but the fact that she is or is not pregnant. Most of the signs of pregnancy are only conjectural, and although an obstetrician of great experience might probably form a fairly accurate forecast in the absence of the really diagnostic signs he would hesitate to make a positive statement on oath unless the infallible signs were present.

The two signs of pregnancy usually depended on by married women to determine their condition are the temporary cessation of the menses and the sensation of quickening. Neither of these signs is of value to the medical jurist, seeing that the woman he interrogates is an unfriendly witness. It is usually her object to impress the medical examiner with the idea that she is pregnant (in some cases the converse obtains), and therefore no reliance can be placed on her statements in this respect. If a truthful statement as to these signs is given, it is of little value, since the stoppage of menstruation may be due to other causes than pregnancy, and the sensation of quickening is a subjective indication, and is unreliable even in the case of women who have experienced it in former pregnancies. In the earliest months of pregnancy the **breasts** enlarge and become firmer. They are more sensitive to the touch, and knotty cords can be felt in the region of the nipples, the areolæ surrounding the nipples appear swollen and shiny and acquire a darker hue from increase of pigment, especially in dark-skinned women. A number of elevated nodules form round the periphery of the areolæ indicating the presence of active sebaceous glands. When the breasts are gently squeezed a milk-like secretion exudes from the nipples. The **abdomen** begins to show indications of distension about the third month, the uterus at this period lies in the hypogastrium. During the first three months the organ retains its pear-shaped outline, afterwards it develops laterally and becomes more ovoid. If the abdominal walls permit of an accurate examination of the shape of the uterus, its form when gravid, together with its mobility and elasticity to the touch, enables the experienced practitioner to arrive at a fairly accurate diagnosis during the earlier months. At, or soon after, this period, alternate contraction and relaxation of the uterine walls may be detected by palpation. The vaginal portion of the **cervix becomes soft** and yielding from infiltration with serum, the edges of the os being rounded, owing to the general tumefaction of the surrounding parts the cervix seems shortened. The mucous membrane of the vagina acquires a cyanotic appearance, and the small veins of the vulva are enlarged and prominent. Bimanual examination—one hand grasping the fundus, and the first and middle fingers of the other applied within the vagina to the os—enables the operator to ascertain the size of the uterus, and probably, in the earlier months—by means of reciprocal movements of the two hands—to feel the sensation of a mobile body within it. The **uterine souffle** is a blowing sound synchronous with the maternal pulse, it chiefly arises in the arteries which run up the cervix, but extends to various parts of the uterus, it may be heard about the fourth month or even earlier.

The signs so far enumerated are suggestive, but not conclusive. There are but *two incontrovertible signs* of pregnancy—the **sounds of the foetal heart**, and the presence and movement of the **foetal members** as felt through the abdominal walls. The foetal heart beats at the rate of 120 to 160 in the minute, the sounds



resemble the tick of a watch at some distance from the ear. When the fœtus is in the usual position they are best heard to the left of the umbilicus, a little below it. They become audible about the 18th or 20th week, but at times they may be heard earlier. If the child is surrounded by an unusually large amount of liquor amni, or if there is a very thick layer of fat on the maternal abdomen, the fœtal heart-sounds are heard with difficulty if at all. By careful palpation the parts of the child's body that are accessible through the abdominal parietes of the mother may be distinguished, and in the living child movements may be felt. Here also the conditions which impede recognition of the fœtal heart-sounds will probably prevent manipulation of the fœtal members, unless the contour of the child's body can be detected, the perception of movements cannot be relied on as a proof of pregnancy, since erratic contraction of the recti muscles of the maternal abdomen might be mistaken for them. If the child is dead the recognition of the fœtal members is the only positive sign of pregnancy.

Since the infallible signs of pregnancy are not available during the first eighteen weeks of pregnancy, no positive statement on oath can be made until this time has expired, which is equivalent to saying that the occurrence of pregnancy cannot be positively determined until quickening has taken place. A negative result yielded by an examination made after a supposed pregnancy of eighteen weeks' duration affords no proof that the woman is not pregnant, the difficulties already mentioned may prevent recognition of the signs of pregnancy, and a further difficulty may be interposed by the child being small and of feeble vitality. As in all other propositions, it is easier to prove a positive than a negative, if the fœtal heart-sounds are heard, whether the body can be felt or not, the fact of pregnancy is established, if neither one nor the other is discoverable, no positive conclusion for the moment can be arrived at. It is under conditions like these that the suggestive signs of pregnancy are of value, as affording grounds for delay and subsequent re-examination or not, as the case may be. Usually a woman pleading pregnancy will state that she is several months advanced, so that some indications of the alleged condition will be present, should she state that she has been only recently impregnated, further examination must be postponed whether there are any indications of pregnancy or not.

### POST-MORTEM APPEARANCES OF PREGNANCY.

Little need be said under this head. In addition to many of the objective signs of pregnancy in the living, two further indications are described, of which the first only is of importance. (a) The presence of an **ovum with villi**, or of a **fœtus** with placental attachment. (b) The presence of a so-called **true corpus luteum** in one of the ovaries. If an impregnated ovum, sufficiently developed as to be recognised as such, is found in the uterus after death (or, in case of ectopic gestation, elsewhere), the fact of pregnancy is established. Certain abnormal products of conception may replace the ovum or fœtus, it is, therefore, necessary to distinguish between such abnormal products and other pathological conditions which occur independently of impregnation. These abnormal products of conception are called **moles**. They are of two kinds—the *sanguineous mole* and the *vesicular mole*.

The **sanguineous mole** is the result of hæmorrhage into the fœtal membranes. The embryo perishes in consequence of pressure produced either by bulging of the chorion and amnion into the fœtal cavity, or the chorion is ruptured and the fœtal cavity is distended with blood, the embryo undergoes maceration

and disappears, or it escapes if the ovum ruptures. If the membranes with the blood-clots remain long in the uterus, calcareous deposits may occur in them. In some cases the distended chorion and amnion form what are known as blood-cysts. The sanguineous mole rarely exceeds the size of an orange. A mole may exist along with a foetus which undergoes normal development, twin conception having taken place. One of the ova develops, the other degenerates.

The **vesicular mole** is formed by vesicular degeneration of the villi of the chorion. The appearance is that of a vast number of vesicles varying from the size of a pin's head to that of a pigeon's egg. The vesicles are, as it were, threaded in rows, cysts forming along the length of the villi, the cysts are filled with a fluid containing mucin and albumin in varying proportions. The embryo perishes, and if the degenerative process commences at an early period of pregnancy all traces of it may have disappeared when the mole is examined, if it commences at a later period remains of the embryo or foetus may be found.

It is unfortunate that the term 'hydatid mole' has been applied to the results of vesicular degeneration of the chorionic villi, a condition which has no relation with the true hydatid. The distinction involves more than mere pathological accuracy—it determines the important question—Can a "hydatid mole" be formed without impregnation? From the description given, it will be seen that the so-called "hydatid" or vesicular mole results from degeneration of a product of conception, therefore it can only occur as the result of impregnation. The true hydatid (*echinococcus*) is very rarely found in the uterus. Gunsburg<sup>1</sup> reports a case in which labour was impeded by a true hydatid growing from the lower part of the uterus, and states that he found only four cases recorded of hydatid tumour in the cavity of the pelvis.

**Corpus Luteum.**—The difference between the so-called *true* and *false* corpora lutea was formerly regarded as, in itself, constituting proof of the pregnant or non-pregnant condition having obtained during life.

The **false corpus luteum**, or as it is also called, the corpus luteum of menstruation, does not undergo development under the usual conditions.

The **true corpus luteum**, or corpus luteum of pregnancy, develops for several months after impregnation, and attains a very much larger size. This increased development is due to greater activity of the circulation, and probably also to trophic nerve influences resulting from the presence of the ovum in the uterus. Such increased development, however, may take place in consequence of myomata and other pathological conditions of the uterus. On the other hand, pregnancy may occur without the formation of a true corpus luteum. It may be accepted for medico-legal purposes that corpora lutea, as indications of pregnancy or of non-pregnancy, are of no diagnostic value.

## SIGNS OF DELIVERY.

The question of recent delivery most frequently comes under the notice of the medical jurist in relation to infanticide or to concealment of birth. In civil cases feigned delivery may require investigation. In respect to imputations against the chastity of unmarried women and under other circumstances, an opinion as to the occurrence of remote delivery may be required.

Taking **recent delivery** first, the subject is divisible into the signs observable in the **living**, and those observable in the **dead**.

<sup>1</sup> *Centralblatt für Gynäkologie*, 1884.

**SIGNS IN THE LIVING OF RECENT DELIVERY AT TERM.**

A woman who has been delivered at term within two or three days of the period of examination presents more or less the following appearances. There is usually a certain languid look, like that of a person recovering from an illness. The lower eyelid and its surroundings are pigmented to an extent which varies with the complexion of the individual. The **temperature** may be slightly elevated. The **pulse** slows immediately after labour, and then quickens, and again slows, and remains so for several days, it is full, but there is no increase in arterial tension. The **skin** is moist. The **breasts** are full and elastic, or they may be hard and nodulated, the superficial veins are visible. The nipples are described in the pregnant condition. The fluid **colostrum** - at first secreted is not true milk, it is viscid, and contain yellow particles visible to the naked eye. Microscopical examination shows the presence of large corpuscles called **Colostrum corpuscles**, which are composed of a number of fat granules bound together by a hyaline substance. According to Heidenhain, these corpuscles are cells of alveolar epithelium, which become round and faintly granular, and eventually take up fat granules from the alveoli, when treated with acetic acid they display nuclei. True milk contains very few colostrum corpuscles, and less albumin than is present in colostrum. The **abdominal walls** are flaccid and wrinkled, beneath them the **uterus** is felt as a hard round ball. The **cervix** is soft, and open at the internal os, the external os is patulous, the lips being bruised, and they may be torn. The internal os begins to close during the first twenty-four hours, the external os remains patulous for a long time. The **vagina** is dilated and relaxed, the rugæ being absent. The mucous membrane at the mouth of the vagina usually exhibits some slight tears, especially in primiparæ, in them also any remains of the hymen are completely destroyed, as evinced by the presence of recent tumefied nodules (*caruncula myrtiliformes*). The **vulva** is tumid and open, especially at its perineal aspect. The **posterior commissure** is usually ruptured, the perineum in some cases being lacerated. The **lochia**, at first almost pure blood, change about the third day to a serous fluid more or less tinged with blood, containing epithelium, mucus, exudation-corpuscles, and shreds of membrane. Subsequently, the blood diminishes, and its place is taken by fatty granules and pus, the colour changing to yellowish or greenish. The amount and the duration of the lochia are variable. The signs of recent delivery are well-marked for the first few days, after the lapse of a week they are more difficult of recognition, and in a fortnight will have so far disappeared as to render absolute diagnosis of *recent* delivery impossible. It is to be understood that the signs enumerated refer to those which follow delivery at term.

**SIGNS IN THE DEAD OF RECENT DELIVERY AT TERM.**

In addition to the indications available in the living, ocular inspection of the uterus and its appendages becomes feasible when a necropsy is made of the body of a woman recently delivered. If death has taken place soon after delivery, before the commencement of involution, the **uterus** will present the appearance of a flabby bag nine or ten inches long, with widely open mouth. Within, the surface is irregular and is covered with coagula of blood, with portions of decidua and with flakes of lymph. At the part where the placenta was attached the muscular structure is devoid of covering, and is darker in colour than the rest of the organ. Here may be seen as large lacunæ the openings of the dilated veins (*uterine sinuses*). The **cervix** is drawn out and is much thinner than the

walls of the body of the uterus, it often presents a bruised and ecchymosed appearance. The **vagina** is dilated, relaxed, and devoid of rugæ. A so-called true corpus luteum will probably be found in one of the ovaries.

### SIGNS IN THE LIVING OF REMOTE DELIVERY AT TERM.

In a woman who has given birth to a child at some remote period the **breasts** are more pendulous and flaccid, and the nipples usually more prominent, the areolæ being deeper in tint than in nulliparous women. The **abdominal walls** also are not so firm and elastic, and the skin of the abdomen is usually marked with **streaks** of a silvery lustre, these marks are not invariably found after pregnancy, they are wanting in about 8 per cent. of cases that have gone on to full term. They occur from other causes than pregnancy from peritoneal tumours, ascites, ovarian tumours, and on the legs from typhoid and typhus fevers, in pregnant women they also occur on the breasts. Although often called cicatrices they do not result from replacement of one tissue by another, which is the characteristic of a cicatrix. They are due to partial absorption of some of the elements of the skin and to modifications of others. The fibrous tissue of the corium, which is normally arranged in a kind of dense network, is rearranged in parallel lines which traverse the streaks from side to side, the papillæ are diminished in size by atrophy, and are spread wider apart. The **labia** are more open and the **posterior commissure** is usually represented by a cicatrix which may extend to the perineum. The **hymen** will not only be replaced by carunculæ myrtiformes, but the continuousness of the nodules composing the carunculæ will be destroyed. The formation of carunculæ results from coitus, but the base of the hymen remains where the mucous membrane of the vagina is folded over to form it and is only completely destroyed by the passage of the child's head. The **vagina** is more open and smoother than before child-bearing, its anterior wall may project into the lumen of the canal. The **cervix** uteri is irregular, the os is puckered, fissured, and more circular and patent than in nulliparous women, admitting the tip of the finger.

These signs are best marked in women who have borne many children. When a woman has borne one child at a period remote from the time of examination, many of the signs may be wanting. The most deceptive appearance is presented by women who have given birth to a single child short of the full term. In such cases, after the lapse of a few years there may be no indication that the woman has been delivered even two or three deliveries of this kind may leave no trace, the woman presenting all the characteristics of a nullipara.

### SIGNS IN THE DEAD OF REMOTE DELIVERY AT TERM.

The only additional evidence to that afforded by the signs of delivery in the living is obtained from inspection of the walls and cavity of the uterus. The uterus that has contained a child remains permanently larger and heavier than in the virgin state. The walls are thicker, and the cavity is not so triangular in outline, the angles where the Fallopian tubes enter being rounded off. Evidence of past rents of the external os may be seen as cicatricial irregularities. The difference between a parous and a nulliparous uterus, however, may not be sufficiently well marked in a given case to enable a decision to be arrived at.<sup>1</sup>

<sup>1</sup> See *Trans. of the Obstetrical Society*, vols. xvii and xviii.

## CHAPTER XIV

### CRIMINAL ABORTION.

IN medical language the term abortion refers to the expulsion of a fœtus, or an embryo, before the viable period — *i.e.*, before the twenty-eight week of gestation. Delivery after the twenty-eighth week, but before full term, is called premature labour. The word miscarriage is used for any interruption of pregnancy before term. Under the term abortion, the law includes both these periods, therefore criminal abortion consists in unlawfully procuring the expulsion of the contents of the gravid uterus at *any period of gestation short of full term*. It will be observed that this definition does not provide for the induction of premature labour by medical men. The law does not recognise this proceeding by making any exception in its favour, hence the necessity for medical men to protect themselves when about to induce labour prematurely, for the purpose of saving the mother's life, or from other proper motives, by explaining the object and necessity of the operation to all concerned, and, if possible, obtaining the moral support of a colleague. Under certain conditions the necessity for the operation is universally admitted, and if the practitioner under such conditions acts openly, he is held to be exonerated from the penalties imposed by the law, this does not mean that legal proceedings cannot be instituted, but no conviction would ensue.

The law is thus expressed —

“ Every woman being with child, who, with intent to procure her own miscarriage, shall unlawfully administer to herself any poison or other noxious thing, or shall unlawfully use any instrument or other means whatsoever, with like intent, and whosoever with intent to procure the miscarriage of any woman, whether she be or be not with child, shall unlawfully administer to her, or cause to be taken by her, any poison or other noxious thing, or shall unlawfully use any instrument or other means whatsoever with like intent, shall be guilty of felony ”

“ Whosoever shall unlawfully supply or procure any poison or other noxious thing, or any instrument or thing whatsoever, knowing that the same is intended to be unlawfully used or employed with the intent to procure the miscarriage of any woman whether she be or be not with child, shall be guilty of a misdemeanour ”  
(24 and 25 Vict., ch. 100, ss. 58 and 59 )

Attention is first directed to the statement that the intention of the act constitutes the crime. If means are taken to produce abortion it matters not whether the woman is or is not pregnant, and further, if the woman is pregnant, it is not necessary that abortion should follow the attempt to procure it in order to constitute the crime. If a woman places herself in the hands of any one with the object of having abortion criminally induced, and she dies in consequence, the operator or administrator is guilty of murder, although he or she had no intention of causing the woman's death. If the means employed were of such a nature as not to be dangerous to life, the crime may be reduced to manslaughter. This throws great responsibility on medical witnesses. In the first of the two sections of the Act quoted, it is stated that the administration of “ any poison or other noxious thing ” with intent to procure abortion is felony. The question becomes what is a noxious thing? Any substance, however

harmless, under certain conditions or in certain amount may be held to be injurious, much depends on the intent with which it is administered. For example, if a large dose of jalap is secretly administered as a practical joke, or in some alcoholic beverage, with the object of punishing a surreptitious drinker of the same, the law regards such a proceeding as a venial offence. If the same drug is administered to a pregnant woman, or to a woman whom the administrator believes to be pregnant, with the object of procuring abortion, it would probably be regarded as a 'noxious thing', not because the administrator is of necessity likely to be followed by abortion, but on account of the criminal intention it displays (see Section 58 above quoted). It might be supposed that the word "noxious" was redundant in relation to the phrase "other means whatsoever," inasmuch as the latter covers all the ground, it is not improbable, however, that the framers of the Act had in his mind a reference to mechanical or local means of procuring abortion. The first clause specifies poison, and then strengthens or enlarges the term by adding 'or other noxious thing'. The second clause specifies "any instrument," and to prevent quibbling adds "or other means whatsoever". One judge, however (Brett), ruled in a case of criminal abortion, that the word "noxious" should be omitted in the indictment, as the comprehensive phrase "other means whatsoever" included any substance administered whether noxious or not (*Reg v Willis*, 1871). As this ruling does not universally obtain, it is necessary for the medical witness to be prepared to express an opinion in the witness-box as to the properties, noxious or otherwise, of the drugs most commonly used with the object of procuring criminal abortion.

### MODES OF PROCURING CRIMINAL ABORTION.

Criminal abortion is attempted in one or more of three ways—By the *administration of drugs* by the mouth, by acts of *general violence*, by *mechanical means* used directly to the uterus or its contents.

Two classes of **drugs** which are reputed to act directly on the uterus include many of the substances used for procuring abortion criminally—they are *emmenagogues* and *ecbolics*. The former are supposed to promote menstruation or to re-establish it after its arrest from causes other than pregnancy, the latter are supposed to cause the expulsion of the contents of the gravid uterus. Whatever value this distinction may have in therapeutics, it is not without importance to the medical jurist. In the case of those who have even a limited acquaintance with the properties of drugs, the administration of a drug supposed to possess ecboic properties would have much more significance than if it was known merely as an emmenagogue. The accused may state that he did not know the woman was pregnant, he was only told that there was a temporary absence of menstruation, the administration of an emmenagogue would be consistent with such a statement, but not that of a reputed ecboic, such a drug would only be given when the object of the administrator was to empty a gravid uterus. Unfortunately the distinction between these two classes of drugs is not well defined, many drugs are included in both. There are, however, two or three that would not be administered for simple stoppage of menstruation.

A favourite emmenagogue of the working classes is **pennyroyal** (*Mentha pulegium*), usually given as pennyroyal tea, it has also been used for procuring abortion, and is about as efficacious in one direction as in the other. This is a substance that might be innocently administered to a woman in the early stage or pregnancy, the administrator regarding the case as one of simple arrest

of menstruation *Savin* (*Juniperus sabina*) is another so-called emmenagogue, but with a different reputation, it is better known as an abortive. The selection of this drug for the purpose of restoring arrested menstruation would be suspicious of criminal intent. Savin is an irritant, and possesses no real emmenagogue nor ecboic properties. The like may be said of a number of substances popularly regarded as abortifacients, as rue, tansy, yew, saffron, etc. In addition to reputed ecboics, powerful purgatives, as colocynth, aloes, and gamboge, emetics, as tartar emetic, irritants, as arsenic, salts of lead, phosphorus, cantharides, and hellebore, have been resorted to. Quinine in large doses (20 grs) causes contraction of the gravid uterus, and has been taken improperly. Pituitary extract is also a powerful stimulant to uterine contractions.

The drug which is most often regarded as an ecboic is *ergot* (*Secale cornutum*). When labour at term has commenced, the contractions of the uterus can both be increased and altered in character—becoming more tetanic—by the administration of ergot. This specific action on the uterus has led to the use of the drug for this purpose of initiating contraction of the gravid but quiescent organ. Experiments on animals justify, but experience of its use with the human female negatives, the views held by many as to its efficiency in procuring abortion. Robert<sup>1</sup> states that two constituents of ergot, cornutine and sphacelmic acid, produce abortion in dogs and cats. In the earlier months of pregnancy (the period when abortion is resorted to) the administration of ergot will not induce miscarriage in the quiescent human uterus, except the organ is predisposed to part with its contents on account of a diseased state of the membranes of the ovum, such as occurs in syphilitic subjects, or from general disturbance of the system caused by the toxic action of the drug. Even when abortion is threatened—as evinced by hæmorrhage—the action of ergot does not necessarily stimulate the uterus to expulsive efforts. Atthill<sup>2</sup> gave it in such a case with the object of emptying the uterus, the hæmorrhage was arrested, but no abortion resulted, the patient gave birth in due time to a living child. Even when full term is approaching, ergot will rarely initiate labour. Arrhill<sup>3</sup> states that he has frequently given ergot as a preventive of post-partem hæmorrhage, commencing its administration a week or ten days before the expected advent of labour, and never once had reason to suppose that it hastened that event. Saxinger<sup>4</sup> records a case in which a woman with contracted pelvis, on two occasions had premature labour induced by means of the introduction of a bougie. On the third occasion—four or five weeks before term—powdered ergot was tried in place of mechanical means. Four grms (62 grains) of the fresh powder were given daily for three days, when the pulse fell from 88 to 60 beats per minute, and the patient was attacked with vomiting and feebleness, but the uterus did not respond. The experience of many other obstetricians corroborates the conclusion that, in the absence of predisposing conditions, ergot in non-poisonous doses has not the power of developing uterine action in the quiescent gravid state.

When abortion has resulted from the use of ergot it has been due to the general toxic effects produced. Richter<sup>5</sup> reports the case of a girl six to seven months pregnant, who took from 2 to 4 ounces. She suffered from the symptoms of acute ergot-poisoning—quick pulse, thirst, pain in stomach and abdomen, stoppage of urine, great restlessness—and, half an hour after giving birth to

<sup>1</sup> *The Practitioner*, 1885

<sup>2</sup> *Dublin Journ. of Med. Science*, 1888

<sup>3</sup> *Brit. Med. Journ.*, 1889

<sup>4</sup> *Maschka's Handbuch*, Bd. 3

<sup>5</sup> *Vertheiljahreschr. f. ger. Med.*, 1861

a dead child, died from profuse hæmorrhage Tardieu<sup>1</sup> reports the case of a woman four months pregnant who aborted in consequence of taking ergot, and died from peritonitis twenty-four hours afterwards In another case, reported by Otto,<sup>2</sup> death took place shortly after the expulsion of an embryo five inches in length Death may result from ergot without abortion having occurred, illustrations of this will be found in the toxicological section on ergot poisoning

During recent years the use of *lead*, most frequently in the form of pills made from diachylon, has become fairly common The practice appears to have originated in the Potteries, where women working with lead would be likely to discover its effects in this direction Hall<sup>3</sup> who has made an exhaustive enquiry into the subject, states that there have been hundreds of cases in the Midlands, and that from that area its use has spread to various parts of England, while cases have also been recorded in Scotland and Wales The information appears to be handed on by women of the midwife class, and there is also a belief that taking lead before menstruation will shorten the period and prevent conception occurring Lead undoubtedly has a powerful effect in bringing about expulsion of the uterine contents, but it is not likely to do this without producing the severer symptoms of plumbism It is probable that many of the women who take diachylon are unaware of its poisonous action, or even, outside the Potteries, that it contains lead Unexplained cases of plumbism in pregnant women should now always be looked upon with suspicion

It may be inferred, from a consideration of the cases recorded in which abortion has been attempted by resort to drugs, that no infallible *cebole* is known When abortion has resulted from the uses of drugs, it has been due either to a predisposing condition of the uterus or of its contents, or to the general toxic effects of the substance taken

**General Violence as a Cause of Criminal Abortion.**—The universal experience of medical practitioners is that under certain conditions pregnant women may suffer from general mechanical violence of an extreme kind without causing miscarriage, and under other conditions the uterus rids itself of its contents on the slightest provocation Since excessive violence to one woman fails to produce results which follow the occurrence of a mere shadow of it to another, it is obvious that the result is determined by intrinsic rather than by extrinsic relations So long as the gravid uterus and its contents maintain their normal healthy relations towards each other, general violence is not usually successful unless the mother's life is endangered and not always then

When general violence is resorted to with the intention of causing abortion, one or more of the following methods may be adopted —Purposely falling down stairs, or from a height, the use of excessively tight stays, allowing the abdomen to be violently kneaded, walking, or riding on horseback, for many successive hours until the bodily powers are quite exhausted, and other means, all of which are singularly futile Such are usually the prelude to more direct and efficacious procedures

**Local Violence as a Cause of Criminal Abortion.**—It has been shown that drugs and general violence, as a rule, fail to bring about miscarriage Greater success attends the use of mechanical means, applied in such a way as to disturb the relation between the uterus and its contents To accomplish this the force must be applied directly to the parts in question, the usual method is to pass up the vagina, in the direction of the womb, some pointed instrument, such as

<sup>1</sup> *Annales d'Hygiène*, 1855

<sup>2</sup> *Maschka's Handbuch*, Bd 3

<sup>3</sup> International Medical Congress, 1913



a piece of wire or of wood, the object being to perforate the membranes through the os uteri. The operation is of ancient date, it is described by Soranus of Ephesus, who flourished about the end of the first century. The direction actually taken by the instrument depends upon the knowledge and skill of the operator. An abortionist who has some slight acquaintance with the anatomy of the parts, and has had sufficient practical experience to enable him to utilise it, will at least endeavour to pass the implement through the os uteri, with the object of puncturing the membranes of the ovum. A less informed person, who has probably only heard the method described as "passing a piece of wire up the private parts till blood comes," will perforate the cervix, or the body of the uterus, in the endeavour to reach its contents. Unqualified practitioners of a low type, and some who are qualified, use male metallic catheters, and by taking care not to wound the uterus, are not unfrequently successful in accomplishing their object without much disturbance of the woman's health. There are reasons for believing that abortion is criminally procured in a vast number of cases in which the operation is undertaken by men who have, or who have had, some relation with the medical profession, and who are, therefore, capable of exercising such a degree of skill in puncturing the membranes as to avoid injuring the uterus, thus materially diminishing the risk of the operation, and consequently lessens the probability of the crime being detected. It appears to be the practice of some abortionists after passing an instrument to advise the woman to consult a registered practitioner as soon as there is any pain or hæmorrhage. The case thus comes before him as a natural abortion, and suspicion is less likely to be aroused.

Cases in which criminal abortion has been procured rarely come before a court of law except when the result of the proceeding has been fatal to the mother. As the woman solicits the crime, and is a culpable party, it is only natural that she should strive to conceal it, in this she is usually successful unless her life is endangered, when the necessity for legitimate medical advice puts others in possession of the secret. Fatal results are mostly due to ignorance or to recklessness on the part of the operator, and to carelessness on the part of the patient. It is well known to obstetricians that puncturing the foetal membranes is not always followed by immediate expulsion of the uterine contents, days of inaction may intervene. Under such circumstances the woman becomes impatient, and urges the abortionist to take more active measures. In this way even a practised hand may be induced to resort to unnecessary violence by which he perforates or lacerates the uterine mucous membrane, and so gives rise to septicæmia. Abortionists are not unfrequently of intemperate habits, and under the influence of drink they lose their usual cautiousness, and do more damage than they intended. The patient also being obliged to comport herself as though nothing was the matter, goes about her daily work, and thus adds another element of risk.

The procurers of criminal abortion sometimes adopt more heroic means than simple puncture of the membranes. Wherry<sup>1</sup> records the case of a woman three months pregnant who had the foetus cut in pieces *in utero* by an abortionist with what she described as a "silver hook." The legs, the arms, and the head were separated from the trunk, the fragments coming away piecemeal, they showed no signs of decomposition, the skin being rosy and firm. Peritonitis ensued, but the woman recovered.

Attempts are occasionally made by unscrupulous women to avail themselves by fraud of the skill of honourable members of the medical profession in order

<sup>1</sup> *Brit. Med. Journal*, 1881

to procure abortion. Such women will go to the consulting-room of a medical man, or to the out-patients' room of a hospital, and state that they are suffering from displacement of the womb, adding that the present is not the first time that the organ has become displaced, and that on previous occasions the doctor they consulted straightened or replaced it with the aid of the uterine sound. It is needless to say that inquiries as to the possibility of pregnancy are skilfully evaded, and as the fraud is attempted in the earliest months, a medical man off his guard is easily deceived. Miscarriage may be induced by passing a bougie between the ovum and the uterine walls and allowing it to remain until expulsion takes place, by vaginal douches of warm water, and by dilatation of the cervical canal by means of elastic dilators, methods usually only adopted by skilled persons. De la Touche<sup>1</sup> relates the case of a pregnant woman who herself introduced the nozzle of a syringe into the os uteri, a female abortionist then manipulated the ball of the syringe, but before half the injection had reached the uterus the patient fainted and died forthwith. At the necropsy the membranes were found to be detached from the lower segment of the uterus, but the placental zone was intact. Death was apparently due to syncope, as nothing else abnormal was discovered. Syncope, which may be fatal, occasionally occurs after the introduction of the uterine sound. See section on sudden death, p. 31.

Occasionally the **woman herself** attempts to procure her own **miscarriage** by direct mechanical means. Various implements have been used with this object. Partridge<sup>2</sup> relates the case of a woman in the seventh month of pregnancy who introduced a hair-pin, points downward, into the uterus and allowed it to remain. No abortion took place for three weeks, the woman then became ill and sought medical advice, when the hair-pin was found to have penetrated the walls of the uterus. The os was dilated, and the hair-pin and ovum were extracted twenty-three days after the introduction of the former, the patient died of peritonitis. Green<sup>3</sup> describes the case of a married woman who introduced a crochet-hook into her uterus, a plan which she had found successful on a previous occasion. The hook slipped from her fingers and she was unable to recover it, though she sought for it with a "marrow spoon." When seen three days later, she complained of pain in the lower part of the abdomen, and there had been slight hæmorrhage from the vagina. Three days later she said she could feel the hook in the abdomen, and indicated a spot midway between the umbilicus and the pubes, where a hard foreign body was palpable. On making an incision, the hook, which was of bone and  $4\frac{1}{2}$  inches long, was found lying between the layers of the mesentery. The woman made an uninterrupted recovery. It was remarkable that the passage of an unsterilised instrument into the abdomen and its presence there for eleven days caused no septic trouble.

Percy<sup>4</sup> records a case in which a young woman attempted self abortion by introducing a catheter into the vagina. Pain and flow of blood followed, and two days later she consulted a physician, who decided that the abortion should be completed. "The doctor invited instruments and worked for a long time, causing most excruciating pain and severe bleeding." After two hours he encountered a loop of intestine in the vagina, and he then packed the vagina and sent the patient to hospital. On admission the patient was collapsed and bleeding from the vagina. Laparotomy was performed, and it was found that 20 inches of the sigmoid flexure had been separated from its mesentery. There was a tear in the bladder  $4\frac{1}{2}$  inches long, and perforation of the anterior vaginal wall. A loop of the sigmoid

<sup>1</sup> *Semaine Gynecol.*, 1896

<sup>2</sup> *New York Med. Journ.*, 1884

<sup>3</sup> *Brit. Med. Journ.*, 1912

<sup>4</sup> *Surgical Clinics of Chicago*, vol. 1, 1917

flexure filled the bladder and protruded into the vagina. The abdominal cavity contained blood clots and three wads of cotton wool half the size of a man's fist. The uterus had not been injured, and gestation was undisturbed.

The loop of the sigmoid which had been stripped of its mesentery was resected, and the rents in the vaginal wall and bladder were repaired. The patient made a good recovery, and pregnancy was not interrupted.

The attempt to procure criminal abortion is usually made in the *earlier months* of pregnancy before the enlargement of the abdomen becomes obvious. Young, inexperienced girls as a rule resort to drugs first, and, after proving their inefficacy, apply to the professional abortionist, in this way the actual abortion may be delayed until the third or fourth month or later. Women of more experience seek mechanical aid as soon as they suspect their condition—after the omission of one menstrual period. It is to be remembered that married women, as well as those who are single, have recourse to forced miscarriage. In the earliest months the attachment of the ovum to the uterus is but feeble, and consequently direct mechanical disturbance is almost certain to cause miscarriage, if effective, the uterus expels its contents in from a few hours to three or four days after the operation. It is quite possible, however, for a uterine sound (or other similar instrument) to be introduced into the cavity of the gravid uterus without causing miscarriage, in such cases the membranes are not ruptured, the blunt sound passes between them and the uterine wall, and produces but slight separation. Many gynaecologists have unwittingly passed a sound into a gravid womb without ill effects. Saxinger<sup>1</sup> relates a case in which a medical man (under the impression that he had to deal with a pathological condition of the organ) on two separate occasions passed a sound into the uterus of a pregnant woman in the early months without interfering with the normal course of gestation.

The induction of criminal abortion imperils the patient's life in two ways - by causing *profuse hæmorrhage*, the result of retention of the placenta or some other product of conception, or by *septic inflammatory processes*. Lesser<sup>2</sup> states that in instrumentally induced abortion the extent of injury to the uterus does not materially influence the length of time the woman survives, septicæmia being always the cause of death. Septicæmia following abortion is a suspicious symptom pointing to mechanical interference as the cause. Little surprise can be felt at the frequency with which septic inflammation follows criminal abortion when the absence of all attempts to render the instruments used aseptic is taken into account. It is not to be forgotten, however, that death of the ovum from pathological causes may determine septic inflammation from absorption of the products of putrefaction. The retention of the placenta, or part of it, after spontaneous abortion also frequently gives rise to septicæmia.

### SIGNS OF ABORTION IN THE LIVING.

The indications that abortion has taken place vary with the period of gestation at which it occurred and the interval that elapses between the event and the examination. If pregnancy is cut short in the first or the second month, even on early examination very little will be found different from that which accompanies an ordinary menstrual period, when a few days have elapsed the parts will have entirely recovered their usual condition. Abortion at three or four months leaves immediate traces behind in the form of a more or less

<sup>1</sup> Maschka's *Handbuch*, Bd 3

<sup>2</sup> *Atlas der ger Med* (zweite Abtheilung), 1891

patulous condition of the vagina, and possibly the vulva may be swollen Spiegelberg<sup>1</sup> describes a funnel-shaped condition of the cervix uteri, which grows narrower from below upwards, as being very characteristic of the recent occurrence of abortion. The condition of the breasts should be investigated. The further pregnancy is advanced so much the more will the signs of abortion resemble those already described as attending delivery at term. As the signs of abortion are less pronounced than those of delivery at term, so do they disappear sooner. The examination of a woman after the alleged occurrence of abortion should be conducted within twenty-four hours, otherwise little information will be yielded by it. Pathological indications in the form of metritis or peritonitis may be present.

### SIGNS OF ABORTION IN THE DEAD.

In making the preliminary incisions and in removing the uterus with the vagina attached, great care must be taken not to injure it in any way. If the vagina is slit open some indications may be afforded as to the best direction in which to dissect the uterus, in order to trace out any punctures in its walls, any such puncture, or laceration, should be carefully measured, and its position and direction ascertained. The length and the breadth of the cavity and the thickness of the uterine walls should be measured. The presence in the substance of the uterus of products of inflammation, the position of any perimetritic inflammation, the presence or absence of mucous membrane in the cavity, and the appearance of its walls are respectively to be noted. If the third month of gestation has been accomplished the site of the placenta will probably be distinguishable. Those who have not previously examined the cavity of a uterus recently delivered in the natural way are liable to draw erroneous conclusions from the appearances presented, more especially when the later months of gestation have been reached. The walls of the cavity are black and irregular—as though bruised—and convey the idea that violence has been resorted to, care must be taken to distinguish between the physiological condition and one that results from mechanical ill-usage. The ovaries should be examined for corpora lutea, not that the information thus obtained is of importance, but because questions on the subject may be asked by counsel. In addition to ascertaining the condition of the generative organs the state of the stomach and the intestines must be investigated for possible indication of irritant poisoning resulting from the administration of one or more of the so-called *ecbolics*. For the same reason the kidneys should also be examined. In exceptional cases a caustic fluid has been injected into the vagina for the purpose of inducing abortion, the condition of its mucous membrane is, therefore, to be observed. Lastly, if an embryo is found in the generative tract its probable age is to be ascertained.

The following is an epitome of the **stages of development of the fœtus** during the first five months of intra-uterine life —

**One Month**—The embryo measures about one third of an inch in a straight line from cephalic to caudal curve, and three quarters along the curve, the ovum is about three quarters of an inch long. The presence of the limbs is indicated. The nasal pits, a cleft indicating the position of the mouth, and two black dots representing the eyes, with the umbilical vesicle and the blood vessels, are present. The amnion is close to the embryo, and is separated from the villous chorion by a clear cavity.

<sup>1</sup> *Lehrbuch der Geburtshilfe*

**Two Months.**—The embryo measures half an inch in a straight line from highest point of cephalic curve to caudal curve, along the curve about an inch. It weighs 60 grains. The nasal and oral openings are separated. The head is becoming distinct from the body. The Sylvian fossa is distinguishable. The primitive kidneys (Wolfian bodies) have almost disappeared, and have become divided into urinary and generative organs. The first centres of ossification have appeared in the lower jaw, the clavicle, ribs, and bodies of the vertebrae. The amnion is in contact with the chorion, the amniotic cavity contains more fluid. The villi of the chorion are especially well developed at one spot. The umbilical vessels passing to the chorion are the only visible remains of the allantois.

**Three Months**—The embryo is  $2\frac{1}{2}$  to  $3\frac{1}{2}$  inches long, and weighs from 300 to 450 grains. The head is separated from the trunk by the neck. The ribs are sufficiently developed as to differentiate the chest and abdomen. The papillary membrane is present. The eyelids and the lips are closed. The teeth begin to form. The fingers and toes can be distinguished, also the rudiments of the nails. The penis and clitoris are of equal length, sexual differentiation is commencing. The chorion has lost most of its villi. The placenta is distinct. The umbilical cord is spiral, about  $2\frac{1}{2}$  inches long, and is inserted into the lower fourth of the linea alba. The decidua vera and reflexa are in contact.

**Four Months**—The foetus is 4 to 6 inches long, and weighs from 2 to 4 ounces. The head equals one fourth of the body length. The mouth is open, the nose, eyes, and ears are distinct. The length of the external ear is from 5.5 to 7.5 mm. The skin is firmer. Hairs (lanugo) are beginning to form. The papillary membrane is quite distinct. The eyelids are closed. The occipital lobe is mapped out. Points of ossification are present in the lower segments of sacrum. The placenta is larger, and weighs about  $2\frac{1}{2}$  ounces. The umbilical cord measures  $7\frac{1}{2}$  inches, is more spiral, and is thicker from the formation of Wharton's jelly, it is inserted above the lower fourth of the linea alba. The sex is distinguishable. The chorion and amnion are in contact. Movements of the limbs have been observed.

**Five Months**—The foetus is 7 to  $10\frac{1}{2}$  inches long, and weighs 10 ounces. The head is still disproportionately large, hair commences to appear on it. Lanugo begins to form along the eyebrows and on the forehead. The skin is red, and its surface is covered with *vernix caseosa*. The eyelids are closed, the papillary membrane is still present. The external ear measures 8 to 12 mm in length. The nails are forming. The temporal and frontal opercula grow during this month, and the Sylvian fossa becomes triangular. The surface of the island of Reil is marked by sulci. Points of ossification are present in pubes and os calcis. Bile stained fluid is in the small intestines. The placenta weighs 6 ounces. The umbilical cord measures 12 inches in length. The movements of the foetus are perceptible to the mother.

The subsequent stages of development are described on p. 19

## CHAPTER XV

### INFANTICIDE.

IN its restricted and technical signification, infanticide means the murder of an infant at the time of, or soon after, its birth. To kill an infant a few weeks old would, etymologically, also be infanticide, but such a case presents a different aspect to the medical jurist, inasmuch as it demands only the same kind of evidence as in ordinary cases of murder in older persons, much more is required from the medical witness in a case of true infanticide. In order that murder can be perpetrated the victim must have had a separate existence, "the killing of a child in the act of birth and before it is fully born is not an offence by the present law, although if injuries are inflicted before birth which cause the child's death after birth, the law of murder applies." In the legal sense a partially-born infant has not a separate existence, although it may breathe

and has consequently acquired the power of maintaining life as a distinct individual. For this reason the medical witness in a case of infanticide has not only to give evidence as to cause of death as in ordinary cases of murder, but has further to endeavour to ascertain if the infant breathed, and whether it was or was not born alive. Since infanticide is usually committed with the object of concealing the occurrence of delivery, the deed is done in secret, and in most instances without the aid of a second person, the burden of proof of live birth therefore, falls on the medical witness. The questions to be considered in cases of infanticide relate to the mother on the one hand and to the child on the other.

As regards the **accused woman** the question is. Has she been delivered of a child recently, or at a period consistent with the time that apparently has elapsed since the birth of the infant? The signs of delivery already discussed afford criteria for the solution of this question.

As regards the **child** the questions relate to 1 Its degree of **maturity** 2 Whether it **has or has not breathed**. 3 Whether it was **born alive**, and if so, the **length of time it lived**. 4 The **cause of death**. 5 The **length of time it has been dead**.

## 1. THE DEGREE OF MATURITY OF THE CHILD.

This is to be determined by a minute investigation of the body with reference to the appropriate indications of age which are described in the section devoted to that subject.

## 2. HAS THE CHILD BREATHED?

An answer to this question has to be sought for by examining the **lungs** with respect to (a) **volume**, (b) **colour**, (c) **consistence**, (d) **specific gravity** (*hydrostatic test*). Corroborative evidence may be afforded by the presence or absence of **air in the stomach and intestines**.

(a) **The Volume of the Lungs before and after Respiration.** When the thorax of a newly-born infant that has **not breathed** is opened the lungs are scarcely visible, the thoracic cavity is chiefly occupied by the heart and thymus gland. If these organs are drawn on one side, or the margins of the opening through the thoracic wall are stretched wide apart, the lungs may be seen lying near the vertebral column, their sharp borders reaching forwards to about one-third of the length of the ribs. The diaphragm is only covered by them at its posterior part. If the child has **fully breathed** the lungs more or less fill the thorax and partly cover the pericardium, the right lung being usually more prominent than the left. The thin sharp margins of the lungs have become rounded, and their under surface covers most of the arch of the diaphragm. If the difference in volume between the lungs of an infant that had breathed and one that had not was invariably as pronounced as above described, there would be little need for further evidence. The descriptions given, however, represent the two extremes absolutely no respiration contrasted with fully established respiration, in the latter case there is little probability of error. When, however, the infant has but feebly or imperfectly breathed, the volume of the lungs may be only slightly different— and possibly not at all—from that which obtains in the foetal condition, in such cases nothing is learnt from an inspection of the lungs as to size.

**The position of the Diaphragm.**—The increase in the volume of the lungs causes them not only to spread forwards but also downwards, the result being that the arch of the **diaphragm** is depressed. Usually in infants that have not breathed the highest part of the arch is on a level with the fourth or fifth rib; after respiration it sinks to the level of the sixth or seventh rib. It is obvious from what has already been said in reference to imperfect respiration that the height of the diaphragm can only be distinctive in well-marked cases. It is, therefore, of little use as a means of determining doubtful cases. Even in cases in which infants have fully respired, the diaphragm has been found nearly as high as before the occurrence of respiration. The presence in the abdomen or the thorax of gases or of fluids due to putrefactive processes alters the curvature of the diaphragm.

(b) **The Colour of the Lungs before and after Respiration.**—Before respiration the colour of the lungs is pale brown, resembling that of the liver, but paler. It has been likened by Casper and Laman to strong chocolate and water, sometimes being lighter, more like chocolate with milk, at the borders the colour is a little brighter. The tint varies with the amount of blood present in the lung. The lobes are indicated by lines of a lighter colour. A marked characteristic of lungs which have not respired is that, with insignificant exceptions, the colour is uniform over the entire surface, the posterior portion may be a little darker than the anterior, but there is no mottling.

**After full respiration** the lungs assume a lighter colour, which partakes of many shades from light red to dark bluish-red. When there is much blood in the lungs, dark bluish-red forms the ground-tone on which spots and patches of bright red are seen, when there is less blood the ground tone is light red, the patches then taking the darker hue. This **marbled or mottled appearance** is very characteristic of the **occurrence of respiration**, it is never found in the **œtal lung**. The shaded patches and spots project slightly above the rest of the lung surface, being formed by the distention of the alveoli with air. The **pleura** of the lung that has breathed is very transparent, and imparts brilliancy to the underlying tints, the **foetal lung**, on the contrary, has a dull non-transparent surface. It is absolutely necessary to observe the colour of the lungs soon after the thorax is opened, as exposure to air quickly alters the tints, making them lighter.

When the infant has **feebly respired** for a short time only, indications in the form of spots or patches of a different colour to the rest of the lung may or may not be present on its surface. They usually appear first on the border of the upper lobe of the right lung, and may be found there when absent from the rest of the surface of both lungs. They appear as red or bluish-red spots on the otherwise uniform brownish tint of those parts of the lung into which air has not entered, the spots are irregularly coloured, having a mottled appearance, and being due to inflated alveoli, afford a sure indication that air has entered the lungs. They may be absent, however, in exceptional cases, even when the child has for a short time survived birth, respiration having been imperfectly carried on.

**Artificial inflation** may cause the surface of the lungs to assume a bright red colour, which, according to Casper, is uniformly distributed over the surface, without any trace of mottling. Taylor and Stevenson mention a case in which Braxton Hicks performed artificial inflation in a still-born infant, the attempt at resuscitation being unsuccessful. On opening the thorax it was found that the air cells in about three-fourths of the lungs had received air, but the colour of the lungs was different from that which obtains after

natural respiration, being of a pale fawn tint. On the other hand, Runge<sup>1</sup> and Obolonsky<sup>2</sup> assert that a mottled appearance may be produced by the method of artificial inflation called Schultze's-swinging.

When death results from hæmorrhage in the newly-born infant, the lungs are pale and reddish-grey, if respiration has taken place, bluish-black marbling is to be seen on the light-coloured ground.

To epitomise a uniform brownish colour of the lungs is indicative of the **fœtal state**; a **mottled or marbled appearance**, of various tints as described, is peculiar to lungs that have **breathed**. The possibility of artificial inflation must be taken into account.

(c) **The Consistence of the Lungs before and after Respiration.**—Before respiration the fœtal lung resembles liver, not only in colour but also in texture, it is compact and firm, and on pressure offers resistance to the finger. After respiration the lung is elastic and yielding, and when compressed between the finger and thumb produces a feeling of crepitation. The difference in consistence between fœtal lungs and those which have respired is due respectively to the absence and to the presence of air in the alveoli; therefore, when respiration has been imperfectly performed, the characteristic feel of a lung that has breathed will be only partially present, or it may be absent entirely, if partially present, some portions of the lung will crepitate, others will not.

Coincident with the establishment of respiration is the commencement of the pulmonary circulation. During fœtal life the blood that reaches the lungs is limited to the amount required for their nutrition, when respiration is established, the whole of the blood in the body periodically passes through the lungs, which consequently contain an amount greatly in excess of that which was present in the fœtal state. If an incision is made into the fœtal lung it cuts like liver, and from the cut surfaces the colour of which is uniform—only here and there a little blood appears when slight pressure is made. Lungs that have breathed are less easily cut, as, on account of their elasticity, the tissues recede before the knife, the stroke of which produces a crepitant sound, the cut surface is irregularly coloured, and if scraped with the blade of the knife, blood-stained froth is obtained.

The increased amount of blood that passes through the lungs after respiration is established adds to their weight, on this fact is founded the so-called **static test**. The average weight of lungs before respiration is stated to be from 450 to 600 grains, and after respiration 960 grains, numerous observations have proved that these figures are far from respectively representing the actual lung-weights. As the lungs of the infant under inspection cannot be weighed both before and after respiration, a reliable average weight would be necessary for the utilisation of the static test, this, however, is unattainable, the lung-weights, both before and after respiration, being so extremely variable. Ploucquet proposed to obviate the difficulty introduced by the varying weight of the lungs in newly-born infants, by assuming a certain proportion between the lung-weight and the total body-weight, but the assumption is fallacious, as no such ratio exists. **The static test in any form is useless as a means of determining the occurrence or not of respiration.**

(d) **The Specific Gravity of the Lungs before and after Respiration.**—Although, after the child has breathed, the lungs are statically heavier, they are specifically lighter, because the additional weight caused by the influx of blood is more than counterbalanced by the air that is now present in the alveoli, on the difference in specific gravity of lungs which have and which have not respired,

<sup>1</sup> *Berliner klin. Wochenschr.*, 1882

<sup>2</sup> *Vierteljahrschr. f. ger. Med.*, 1888



is founded the **hydrostatic test**. Advantage is taken of the fact that the specific gravity of foetal lung-tissue is greater than that of water, whilst the specific gravity of lung-tissue which has respired is less than that of water. If, therefore, lungs which have not respired are placed in water they sink if they have respired they float.

**The Hydrostatic Test is thus Performed.**—The lungs, with the bronchi, as far as their junction at the trachea, and the heart are taken out *en masse* and placed in some water contained in a suitable vessel. They either float or sink. If they **float**, observe whether the bulk of them remains above the water-level or whether they float almost or quite submerged. The heart should then be removed, the bronchi divided, and each lung tied separately, the respective degrees of buoyancy being observed as before. Each lung is now cut into about a dozen pieces, and every piece is tested separately, if they float, they are to be taken out of the water and subjected to firm compression, in order, if possible, to drive out the air or gas that causes them to float. If the air has entered the lungs in the act of natural respiration, no degree of force will expel it unless sufficiently powerful to cause disintegration of the lung-tissue, consequently, on again placing the fragments in water they still float. If the lungs **sink**, it should be noticed whether one sinks or both sink, and whether slowly or rapidly. They are then severally divided into about a dozen pieces, each of which is tested separately. If the lungs, when whole and when cut in pieces, sink, presumptive evidence is thereby afforded that the child has not breathed.

The **possible fallacies** of the hydrostatic test are that the lungs may float (α) from *artificial inflation*, (β) from the presence of the *gases of putrefaction*. That they may sink (γ) from the effects of *disease*, (δ) from *imperfect respiration* (*atelectasis*), or from absolute persistence of the foetal condition, although the child has breathed and lived for some time.

(α) **Artificial Inflation.**—This may be performed in several ways by the direct application of the operator's mouth to that of the child, by forcing air into the lungs through a silver catheter, or other similar instrument passed by the mouth into the trachea, or by the method known as Schultze's-swinging. Inflation of the lungs in a new-born infant is no easy matter, the air being apt to find its way into the stomach rather than into the lungs. When the attempt at resuscitation is not successful, *post-mortem* examination shows that the lungs are rarely more than partially inflated. As already described, the appearance of lungs artificially inflated is stated by some to be very different from that displayed by lungs in which complete natural respiration has taken place. It has also been asserted that a further difference exists—that the air contained by lungs artificially inflated can be expelled by pressure. This is contradicted by experience, and is opposed to physiological experiment. In Braxton Hicks' case of artificial inflation, previously mentioned, no amount of pressure short of destruction of tissue caused the divided pieces to sink. An explanation of these contradictory statements may probably be found in the position occupied by the air artificially introduced. If it arrives in the alveoli without rupturing their walls, there is no reason to suppose that it can be more easily expelled by pressure than air which has been naturally inspired. If, however, the air has not penetrated beyond the lobular bronchi, or (as readily occurs in artificial inflation), it has been propelled with sufficient pressure to rupture the septa between the alveoli, and to obtain access to the sub-pleural connective tissue, there is nothing to prevent its expulsion on compression of the lung fragments.

When artificial inflation is only *partially successful* in distending the lungs, it has been stated that no additional blood enters them, so that, although they

may be more voluminous than in the fœtal state, they are not heavier, and that an incision into the lung-substance does not yield blood, as in the case of a lung that has naturally respired. This view, supported by Casper and other leading experts has of late years been proved to be incorrect. Runge,<sup>1</sup> Sommer,<sup>2</sup> and Obolonsky,<sup>3</sup> have all recorded cases in which, by means of Schultze's-swinging the lungs of still-born children have acquired all the properties of lungs that have partially respired in the natural way. Obolonsky removed by Cæsarean section a nine months' fœtus from the body of a woman who died suddenly, the operation being performed about ten minutes after death. The child was dead, and attempts at resuscitation were made by Schultze's-swinging the attempts were unsuccessful, and on examining the body the thoracic cavity was found almost entirely filled with the lungs. The apices of both lungs, and the edges of the lobes were rose-red in colour, and the rest of the lung-surface was purple-red with rose-coloured spots, presenting a mottled or marbled appearance. The lungs floated on the surface of the water and when divided all the pieces floated except those derived from the posterior aspect. From the edges of an incision in the apices blood-stained froth exuded. Other cases are also recorded by the same writer in which blood-stained froth was obtained by the incision of lungs artificially inflated.

The distinction formerly made between the effects of natural respiration and those produced by artificial inflation is no longer to be received in an absolute sense. In many cases it doubtless holds good, but since exceptions have been proved it is to be admitted that *no known test* will enable an infallible opinion to be expressed as to whether lungs have respired incompletely or have been artificially inflated.

Apart from physical evidence, the occurrence of artificial inflation in cases of suspected infanticide is opposed to reason. In the newly-born infant the performance of artificial respiration is a difficult task for the expert, and without special knowledge it would be barely possible for an ordinary woman to put it into practice. Schultze's method, by which lungs have been inflated to a degree comparable with the effects of imperfect respiration, is little used in this country, and demands expert knowledge and practice for its performance. If the inflation is done by a second person who was in attendance at the labour, evidence of the fact would be forthcoming.

(β) **The Gases of Decomposition.**—The lungs belong to the class of organs which putrefy slowly. The thorax being intact, if there are no external signs of putrefaction, or only those of the early stage, it may be accepted that the lungs will not be influenced by the gases of decomposition so far as the hydrostatic test goes. The earliest indication of putrefaction in the lungs consists in the appearance of small vesicles filled with gas between the lung substance and the pleura, the pressure of the gas lifting the pleura in detached spots. They first appear at the free borders of the lobes, and at the base of the lungs, subsequently, the deeper seated tissues of the lungs are infiltrated with gas. The vesicles under the pleura range from the size of a pin's head to that of a bean, in the early stage of putrefaction they are small and are often found clustered together, or in rows. By pressure of the finger the gas can be displaced, and if small the vesicle disappears as the gas travels under the pleura. If such a vesicle is pricked with a needle so as to allow the gas to escape, the pleura falls flat. Air within the alveoli is not expelled, neither by pressure nor by simple pricking with a needle. It is to be noted that an appearance similar

<sup>1</sup> *Berliner klin. Wochenschr.*, 1882

<sup>2</sup> *Vierteljahrsschr. f. ger. Med.*, 1885

<sup>3</sup> *Vierteljahrsschr. f. ger. Med.*, 1886

to the small vesicles of putrefaction might be caused by rupture of the alveoli from too vigorous attempts at artificial inflation. In the early stage of putrefaction the lungs closely retain their original appearance, so that the colour of the foetal lung may be readily distinguished from that of a lung which has breathed. When a more advanced stage is reached the colour is dark green, sometimes almost black, or dirty brown, the lung-substance is softened, and on section a dirty-red fluid escapes: the odour is then highly offensive. Foetal lungs do not decompose so rapidly as those which have breathed.

When foetal lung-tissue is infiltrated with the gases of decomposition, its specific gravity is lessened, therefore in this state it will float. A piece of lung in this condition compressed by the finger and thumb under water gives off a number of relatively large and irregularly formed air-bubbles which ascend through the water, a piece of fresh lung that has breathed when similarly treated yields a stream of fine, equal-sized bubbles. Pressure drives out the gas from decomposed lungs, so that pieces which float in their original condition sink after compression, this result, however may ensue in the case of lungs which have breathed when the lung-substance is softened by putrefaction. Foetal lungs which have been rendered buoyant by putrefaction spontaneously lose that property in a still more advanced stage of decomposition, and sink as they would before being attacked by putrefaction.

It may be accepted that when the lungs are in an advanced stage of putrefaction, no trustworthy evidence can be yielded by the hydrostatic test, the fact mentioned at the beginning of this section, however, is to be remembered - that the lungs putrefy slowly, therefore an advanced stage of external putrefaction does not preclude the necessity of testing the lungs. Ogston<sup>1</sup> found no putrefactive buoyancy of the lungs from the body of an infant which had been dead five months. Casper examined two new-born infants whose bodies were in an advanced stage of decomposition, the heart and the liver floated from the presence of gas, whilst the lungs sank. If a distinction is to be made, the evidence afforded by the lungs *floating* is quite unreliable when the organs are undergoing putrefaction, that afforded by the lungs *sinking* may be taken into account, provided that an advanced stage of putrefaction has not been reached.

(γ) **The Effects of Disease.**- Pulmonary disease in the first days of life is rare, certain forms- as pneumonia, pleurisy with effusion, and so-called pulmonary apoplexy -are capable of depriving the lungs of its buoyancy after the occurrence of respiration. If the disease began during intra-uterine life, it might either completely or partially prevent the entry of air into the lungs. Death from suffocation may produce such a degree of hyperæmia, or œdema of the lungs, as to cause them to sink in water. In none of these conditions would there be any real difficulty, as the pathological appearances caused by the diseases would be apparent either to the naked eye or with the aid of the microscope. With the exception of excessive hyperæmia produced by suffocation, it is unlikely, but not impossible, that the whole of both lungs would be so affected by disease as to sink when subdivided. If sinking of the lungs is due to consolidation after the occurrence of respiration, any attempt to inflate them by blowing down the trachea will be unsuccessful, but if due to persistence of the foetal condition, inflation can be thus accomplished.

(δ) **Imperfect Respiration.**- It is a remarkable fact that very exceptionally an infant may survive its birth for many hours, during which period the chest may rise and fall as in ordinary respiration, and what is even more remarkable,

<sup>1</sup> *Lectures on Medical Jurisprudence*

the child may cry, and yet after death the lungs are found to have completely retained their foetal condition— the colour, volume, consistence, and specific gravity being respectively the same as in lungs which have not respired. From this total absence of aeration, all degrees of alveolar distension may occur up to that which accompanies fully developed respiration. The term **atelectasis** has been applied to such conditions, the meaning of the word being—imperfect expansion; medical jurists use the word in order to indicate defective performance of a physiological function, it is used by others to indicate an acquired pathological condition—partial consolidation of the lungs. The word is unnecessary and inconvenient. The expression “imperfect expansion” is sufficiently distinctive, and admits of change to “non-expansion” to signify absolute persistence of the foetal condition, whereas atelectasis being itself a comparative term cannot correctly be thus qualified.

Several attempts have been made to explain the anomaly of a child living and breathing for many hours without a trace of the occurrence of respiration being present in the lungs after death. Maschka denies that air enters the lungs at all in such cases, and believes that the passage of air along the trachea and bronchi is sufficient to account for the signs manifested during life. Others accept the theory first propounded by Simon Thomas<sup>1</sup>—that in feeble infants the respiratory movements may gradually subside in such a way that the elasticity of the lung-tissue, at every expiration, drives out more air than is drawn in at the inspirations, in this way the lungs, after having breathed, gradually return to the foetal condition. As a result of experimental investigation Ungar<sup>2</sup> states that after respiration has ceased, the air which has entered the lungs may be entirely absorbed by blood still circulating through them.

When from immaturity or from extreme feebleness a newly-born infant barely exists, its demand for oxygen will be very slight, and it is conceivable that the necessary interchange might take place in the air-tubes without the help of the alveoli. The rising and falling of the chest, as in ordinary respiration, is not inconsistent with retention of the foetal condition of the lungs. Hermann<sup>3</sup> showed experimentally that, on account of the adhesion of the bronchial and alveolar epithelium, the lungs in the foetal condition require more pressure to expand them than those which contain air. If the lungs are once expanded, however feeble the infant may be, there will be some movement of air in them, but if the initial unfolding of the lung-tissue has not already taken place, the movements of the chest-wall may be inadequate to overcome the resistance. It is, however, difficult to understand how sufficient air could be inspired under these conditions to enable the child to cry to the extent that has been recorded on several occasions. The second hypothesis—gradual expulsion of all the air by the elasticity of the lung-tissue, is in accord neither with physiological experiments nor with clinical experience. Hermann has proved that the elastic power of lung-tissue that has once contained air cannot expel the air and restore the lung to the foetal state. Children are frequently born immature and feeble, and die shortly after birth, but the occurrence of lungs in a foetal condition under such circumstances is exceptionally rare. Ungar's hypothesis is not improbable, seeing that in these cases of low vitality the heart may continue to beat some time after the cessation of respiration, in this way it seems possible that the residual air might be entirely absorbed by the blood. For example, Causse<sup>4</sup> records the case of a seven months' infant which survived several days, after death the right lung was found more expanded than the left, and,

<sup>1</sup> *Nederl Tijdschr v Geneesk*, 1864

<sup>2</sup> *Vierteljahrsschr f ger Med*, 1883

<sup>3</sup> *Pflüger's Archiv*, 1879

<sup>4</sup> *Annales d'Hygiène*, 1878

when cut into, yielded dark-coloured blood, but it was not crepitant, both lungs, whole and divided, sank in water. In this case it is probable that modified respiration took place in the right lung, as it was more expanded than the left (which remained foetal), and yielded blood on being cut into, still it sank, whole and divided, when placed in water, showing that after death it contained no air.

Whatever may be the true explanation, the fact must not be lost sight of – that under exceptional conditions an infant may breathe and even cry at intervals, for many hours after its birth, and yet after death its lungs may absolutely resemble those of a still-born child. A less extreme condition, in which a portion of the lung contains air, does not present the same difficulty, as the hydrostatic test is capable of demonstrating the fact. It has been asserted that air which has entered the lungs during imperfect respiration can be driven out by compression of the detached pieces of lung so that they will sink in water. If the air has arrived in the alveoli (as it presumably has) it is difficult to see how this can be. When a piece of lung is compressed the bronchioles are compressed as well as the air-cells, the result being that the air contained in the bronchioles is expelled, but not that in the alveoli, the stronger the pressure the more firmly is the air pent up in the alveoli, unless such force is used as to rupture them.

**The inferences** to be drawn from the results obtained **with the hydrostatic test** are – That if the whole of the lungs **float**, both entire, when cut in pieces, and after the application of pressure proof is afforded that **the child has breathed**, the possibility of artificial inflation being borne in mind. If the lungs **sink**, both whole and divided, the **probability** is that the child has **not respired**, this qualified statement is rendered necessary by the occasional occurrence of an imperfect type of respiration, which leaves the lungs in the foetal condition, though not so conducive to error, absence of buoyancy from disease is also to be remembered. It is obvious that no lung-test can distinguish between lungs that retain the foetal type, notwithstanding the occurrence of a modified form of respiration. the stomach-bowel test may be of service in such cases. Less importance is to be attached to negative evidence in this direction, as no injustice to the individual results. at the most, a guilty person may escape, an innocent one cannot suffer. When putrefactive processes have invaded the lungs, the greatest reticence is to be displayed in accepting the evidence afforded by the hydrostatic test, in such cases it is better to state that the condition of the lungs does not admit of an opinion being given as to whether the child did or did not breathe.

**The presence of Air in the Stomach and Intestines.** Breslau,<sup>1</sup> in the year 1865, drew attention to the fact that the stomach and intestines in still-born infants sink when placed in water, whilst in infants which have survived birth a sufficient amount of air is present within one or both of these viscera respectively to render them buoyant. In 1886, Ungar<sup>2</sup> began to advocate the importance of this distinction, which, meanwhile, had not been practically utilised to any great extent, since then the subject has received much attention, especially in Germany, and numerous observations have been made with the view of ascertaining how far the presence or the absence of air in the stomach and intestines can be depended on as proof of live birth, or of the occurrence of respiration. The advocates of this test have designated it “**the second life-test.**”

The air that is contained in the digestive tract of a newly-born infant finds

<sup>1</sup> *Monatsschr f Geburtskunde*, Bd 25

<sup>2</sup> *Vierteljahrsschr f ger Med*, 1887

its way there in the act of swallowing during the first and subsequent respiratory movements, if the child does not survive the first few respirations, the stomach only is more or less inflated, but if the child breathes for some time, the air finds its way into the duodenum and subsequently lower down the intestinal canal. The test is performed by placing double ligatures at the cardiac and pyloric ends of the stomach, and also at the lower part of the duodenum. The stomach, separated from the œsophagus, is with the intestine, then placed in water, if the united viscera float, they are detached and tested separately. The results of many experiments tend to show that this test is of considerable importance in determining the occurrence or absence of respiration. The advocates of the test state that it is not only capable of substantiating the result obtained from the hydrostatic lung-test, but that it is capable of determining whether breathing has or has not occurred in those cases of imperfect respiration in which, after the child has survived several hours, the lungs retain the foetal condition and consequently, yield negative results with the hydrostatic test. There are two ways in which the stomach-bowel test may be rendered untrustworthy: by attempts at artificial inflation, and by the occurrence of putrefaction. Artificial inflation, whether attempted by the direct application of the operator's mouth to that of the child, or by means of a catheter or other tube, or by Schultze's-swinging, is almost certain to cause air to enter the stomach, whether it enters the lungs or not. It is obvious that putrefaction will vitiate the test. In the *absence of artificial inflation* and of *putrefaction*, if the stomach and especially if the duodenum also floats, the fact affords strong evidence that the infant breathed; if the stomach sinks under the same conditions, it affords *no* proof that the child was still-born.

Nikitin<sup>1</sup> gives a table of a hundred cases in which the evidence obtained from the hydrostatic test and the stomach-bowel test are placed side by side. The results show that the latter test is likely to occupy an important position amongst the means resorted to in order to ascertain whether an infant has or has not breathed.

### 3. WAS THE CHILD BORN ALIVE?

It is ever to be borne in mind that the lung test at the most can but prove that the child has or has not breathed. When the medical witness gives evidence based on the lung-test, if the lungs float he must limit himself to the statement that his investigations prove that the child breathed; he must not say that the child was born alive. The reason for this limitation is that, although the written law contains no definition of live birth, the judges do not recognise the birth of a child to be accomplished until the whole of its body is external to the maternal organism, it is not necessary that the umbilical cord should have been divided. As is well known, a child may breathe after its head is extruded from the vagina, the rest of its body remaining within the maternal passages, to kill a child in this stage of its birth is not murder, since, according to the law, it has not then any separate existence. It will be seen from this that proof of complete respiration is not in itself proof of live birth. If the child is killed immediately after its birth, in the legal acceptance of the term, medical evidence cannot prove that it was born alive, no evidence except that of an eye-witness is competent to do so. Again, it is not even necessary for the child's head to be born in order that it may breathe, there are numerous cases on record in which infants have breathed, and have demonstrated the fact by

<sup>1</sup> *Vierteljahrsschr. f. ger. Med.*, 1888

crying whilst still within the uterus (*vagitus uterinus*)<sup>1</sup> McLean<sup>2</sup> records the case of a child that cried for four or five minutes whilst within the uterus, "the voice sounding as if coming from the cellar", delivery was effected with the forceps. Brull<sup>3</sup> delivered a woman of twins after a midwife had vainly endeavoured to do so, the uterus was relaxed and the breech of the first child was impacted in the bum, the child had been heard crying for three hours, and it continued to do so during extraction. In the recorded cases in which respiration took place whilst the child was inside the maternal passages, the admission of air resulted from their dilatation either by the hand of the accoucher or the instruments used by him, so that a channel of communication was established between the atmosphere and the child's mouth. If this is invariably so, the occurrence of respiration whilst the child is within the vagina or uterus would be of no importance in cases of suspected infanticide since the accoucher could give evidence as an eye-witness as to whether the child was or was not born alive. It appears just possible, however, that air might find admission to the child's mouth without adventitious aid.

It is to be noted in any of these modes in which respiration occurs before legal birth that, if death takes place immediately afterwards, the expansion of the lungs will probably be incomplete. For the reasons already given the initial expansion of the lungs is a gradual process, the adhesiveness of the contiguous layers of epithelium in the bronchioles and the alveoli does not yield to the first few inhalations, but the air penetrates step by step until at last the entire respiratory tract is opened up. According to Dohrn,<sup>4</sup> under normal conditions the alveoli are not fully dilated until the second or third day, in weakly children still longer time is required fully to expand the lungs and to remove all trace of their foetal state. If, therefore, the lungs are found completely expanded, it is probable that the child was fully born, as the difficulty of access of air whilst the head is within the vagina, and the probably short delay in the completion of delivery when the head is outside (the body being within), would render the accomplishment of perfect expansion of the lungs unlikely. At the risk of reiteration, attention is directed to the difference in the quality of the evidence afforded by the hydrostatic test as regards proof of respiration and of live birth respectively. If the lungs are fully expanded, so that every separate piece floats after being subjected to pressure, the occurrence of respiration (allowing for the remote contingency of artificial respiration) is proved whilst the same state of the lungs affords only *presumptive* evidence of live birth. In the one case, with due observance of experimental precautions, there can be no doubt as to the truth of the assertion, in the other, it is a surmise, for though highly improbable, it is not impossible for the child to remain with the body unborn (the head being expelled) until the lungs are fully distended.

**The Stomach-bowel Test** affords circumstantial evidence of survival of birth. If the intestines as well as the stomach contain air, and in consequence float when placed on water the absence of attempts at artificial inflation and of putrefaction being understood the probability is that the child did not die immediately after birth, the lower the air permeates the intestinal canal, the greater the probability that the child survived birth.

**The Middle Ear Test.** In the foetal state the middle ear is filled with an embryonic gelatinous mass. Wredin<sup>5</sup> and afterwards Wendt<sup>6</sup> found that when

<sup>1</sup> *Brit Med Journ*, 1912

<sup>2</sup> *American Journal of Obstetrics*, 1889

<sup>3</sup> *Wiener klin Wochenschr*, 1895

<sup>4</sup> *Archiv f Gynaecologie*, 1889

<sup>5</sup> *Otitis media neonatorum*, 1868

<sup>6</sup> *Arch f Heilkunde*, 1873

respiration has fully taken place, this mass is replaced with air. Wendt further states that if attempts at respiration are made whilst the infant is immersed in the amniotic, or in any other fluid, such fluid will be found replacing the gelatinous mass. Ogston,<sup>1</sup> junior, found that the mass disappeared in from a few hours to two or three weeks after birth. Schmaltz<sup>2</sup> states that the mass begins to disappear during foetal life, and that although the influence of respiration on it is recognisable, the absence of the mass is no certain proof that respiration has taken place, he often found the mass in infants that had breathed, and in some instances it was absent in the foetal state. Lesser<sup>3</sup> states that a few respirations produce little effect on the contents of the middle ear, but that they are displaced after several hours' respiration. Lesser also states that in premature children the foetal condition of the middle ear may persist for more than twenty hours after birth.

**Changes in the Umbilical Cord.**—Much difference of opinion exists as to the value of **mummification** of the cord as an indication of live birth. Casper denies that it has any value, Billard and others hold that its occurrence is an indication of extra-uterine life. Under certain conditions it may undoubtedly be of importance as a diagnostic sign. Lowndes<sup>4</sup> relates the case of a new-born child which had probably been dead eleven or twelve days, for half an inch from the navel the cord was perfectly fresh, then came the usual line of demarcation, and the remainder of the cord, about  $2\frac{1}{2}$  inches, was completely mummified, there was no appearance of ligature. The inference was that the child had lived at least twenty-four hours. Such a case is rare but it serves to show that mummification of the cord is not to be overlooked in suspected infanticide. **Normal separation** of the cord, with the accompanying signs of reaction—the line of capillary congestion and the inflammatory exudation-products—are unmistakable indications that the child has survived birth, unfortunately such signs are not available until, at the least, four or five days after birth. Care must be taken not to confound a red line which at birth surrounds the spot where the cord is inserted, with the line of separation. The former is simply a coloured ring without any swelling or indications of inflammation, the latter is a true reactionary area, of a distinctly inflammatory character. If the body of an infant in an advanced stage of putrefaction is found with no remains of the umbilical cord attached, it is not, therefore, to be assumed that natural separation has taken place, the cord may have been detached close to the abdomen by violence, either during life or after death, and the signs which ordinarily serve to differentiate between normal and violent separation will be destroyed by the putrefactive processes.

**The alterations** which take place in the **skin** of an infant after birth, described on p. 20, may be utilised as indications of survival of birth.

The changes which take place in the heart and vessels after the cessation of the foetal circulation are of no value in determining whether the child was born alive or was still-born. Elasser<sup>5</sup> by careful examination proved that the processes by which the foramen ovale, the ductus venosus, and ductus arteriosus are closed, do not commence for several days after birth, they take weeks to accomplish, and they do not follow any regular order.

Occasionally the presence of **milk** or other nutrient substance in the **stomach** may be discovered, and survival of birth thus proved, anything found in the stomach should be submitted to microscopical examination. Fluids in the

<sup>1</sup> *Brit and For Med Chirurg Review*, 1875.

<sup>2</sup> *Arch f Heilkunde*, 1887.

<sup>3</sup> *Vierteljahrschr f ger Med*, 1879.

<sup>4</sup> *Liverpool Med and Chir Journ*, 1889.

<sup>5</sup> *Henke's Zeitschr*, Bd 64.



stomach derived from either the maternal or the foetal organism—as blood, or meconium—prove nothing except that the infant was living about the time of birth. The absence of meconium from the large intestines, though not affording proof of survival after birth, is suggestive of it, the presence of meconium, however, does not indicate still-birth, as frequently it is not expelled for a day or more after birth. The presence or absence of the *Bacillus coli* in the meconium has been held to afford a means of deciding the question of live birth, but, as pointed out by Binda,<sup>1</sup> although the meconium of dead-born children is sterile, so also are the intestinal contents of those born alive, unless they have swallowed food, therefore the presence or absence of *B. coli* merely proves that the child has or has not taken food. In the intestinal contents of infants which had not taken food, Schild<sup>2</sup> exceptionally found a micro-organism resembling the *B. coli* (but not earlier than twenty-two to twenty-four hours after birth) which he thinks is carried down by the saliva that is swallowed.

It is from a consideration of such of the above described signs of live birth as are available that an opinion is to be formed as to whether the child was born alive, and, if so, how long it lived. In some cases there is no difficulty in determining the former point, and but little in approximately estimating the latter, others present features of extreme difficulty, and demand both painstaking investigations and a display of due reserve in the statement of opinions based on such investigations. If the child died immediately after birth it is out of the power of medical evidence to *prove* that it was born alive, whatever reasons there may be for inclining to that opinion. The **reliable signs of survival of birth** only come into play after the lapse of several days.

#### 4. THE CAUSE OF DEATH.

The first consideration is—Did the child die from natural causes? It is a well-ascertained fact that among illegitimate children the mortality is much greater than it is among those born in wedlock. In 1919, the deaths of legitimate infants were 83.7, and of illegitimate infants 172.8 per thousand births in England and Wales. As regards the causes of death, the excess was very slight from infectious diseases in general and from congenital defects, and comparatively so from bronchitis and pneumonia, but heavy for diarrhoea and especially for atrophy. These facts suggest that the illegitimate infant may be at a greater disadvantage in regard to the purity of its food than to protection from the risk of pulmonary infections. The illegitimate mortality is most excessive in the case of deaths attributed to syphilis, being over eight times as heavy. Some of this excess may be due to less reluctance to certify the true cause of death in such cases for illegitimate infants. Fatal injury at birth is also much commoner among the illegitimate, probably owing to many of the confinements taking place under disadvantageous circumstances.

#### DEATH OF THE INFANT FROM NATURAL OR ACCIDENTAL CAUSES.

A large percentage of infants are either still-born or die from natural causes shortly after birth, in England and Wales the proportion of deaths due to premature birth has increased from 11.19 per 1,000 births in 1861, to 18.88 per

<sup>1</sup> *Giornale di Med. Legale*, 1896

<sup>2</sup> *Zeitschr. f. Hygiene u. Inf. krank.*, 1892

1,000 in 1893, and 19.9 in 1919, the deaths from congenital defects have also increased in the same time from 1.76 to 2.63 per 1,000 births in 1919. First pregnancies are more likely to yield still-born children than subsequent pregnancies. In a great many of the cases in which medical examination of the bodies of infants found under suspicious circumstances is required to be made, the mothers are young women who have not previously had children. The necessity for concealing their condition during pregnancy leads to actions which are detrimental to the well-being of the infant *in utero*. The labour also takes place under disadvantageous conditions—usually the woman has no help, and, if parturition is prolonged, or the umbilical cord prolapses, or is coiled round the child's neck, or any other lethal complication capable of being rectified by skilled assistance occurs, the child may be still-born from absence of help, but without any criminal intent on the part of the mother. The infant, from immaturity or intra-uterine malnutrition, may perish during or immediately after birth simply from deficient vitality. Malformations of the heart, of the intestinal tract, or of the central organs of the nervous system are capable of easy recognition as causes of the infant's death. Disease, as apoplexy, pneumonia, hydrothorax, gastro-intestinal hæmorrhage, and the like, may occasion death. Asphyxia, from non-expansion of the lungs or from the introduction of blood, meconium, or liquor amni into the air-passages, is another cause of death.

It will be necessary to describe some of these modes of **natural or accidental death** in detail.

**Prolapse of the umbilical cord** is lethal if the prolapsed part of the cord is subjected to firm and continuous compression sufficient to interrupt the blood current. When the circulation through the cord is arrested, the foetal blood becomes venous and stimulates the respiratory centres into action, the infant makes attempts at respiration and, access of air being prevented, suffocation ensues. The post-mortem **signs of suffocation** will be found to be present on examination of the body—capillary ecchymoses, injection of the tracheal mucous membrane, distention of the heart with blood and the presence of blood, meconium, particles of vernix caseosa, and mucus in the air-passages and stomach, the contents of the bowels are usually voided during the attempts at respiration, and the body of the child is consequently befouled. The discovery of particles of vernix caseosa in the bronchial tubes afford strong evidence in favour of intra-uterine suffocation. Strassmann<sup>1</sup> takes advantage of the fact that weak solutions of aniline dyes stain keratin-tissues. A section of the lung-tissue may be stained in a solution composed of one drop of a 1 per cent solution of gentian-violet to a watch-glass of water, allow to remain five minutes, wash with alcohol, and clear with oil. A simpler way is to take a little of the contents of a bronchial tube, spread it on a cover-glass, dry over a Bunsen flame, and stain, the stain is taken by the particles of vernix caseosa, and their presence is then easily detected.

**Accidental strangulation with the umbilical cord** is frequently stated to have been the cause of death in cases of alleged infanticide, the fact that during birth the child's neck is not unfrequently surrounded by the cord gives colour to the statement. It occurs in about 25 per cent of deliveries, the mortality being estimated at from 1.1 per cent to 2.7 per cent. Sanger<sup>2</sup> relates a case that was delivered in hospital. When the head was expelled, the face was deeply cyanosed, the funis was coiled once round the neck, in the first instance too tightly stretched to be released, and when subsequently removed

<sup>1</sup> *Vierteiljahrsschr. / ger. Med.*, 1887

<sup>2</sup> *Archiv. / Gynecologie*, 1879

no pulsation was perceptible the child was still-born. The skin of the face and neck was of a dark bluish-red colour, sprinkled with numerous petechiæ, very dark almost black, the eye-balls were prominent, the conjunctivæ were swollen and marked with petechiæ, the tongue protruded between the gums, the superficial veins appeared as dark lines. On the front of the neck were two firm ridges, extending to the angle of the lower jaw resembling the "double chin" in stout people, these ridges were covered with large ecchymoses, and between them, over the sternum, was a shallow, blackish groove a few centimetres long. The skin of the trunk was livid and cyanotic. There was a layer of blood 1 cm. thick under the scalp, the brain was anæmic and œdematous, the meninges were pale, and the sinuses only moderately filled with blood. There was fluid in the lateral ventricles. The veins of the skin and of the muscles of the head and neck above the mark of the cord were very hyperæmic, both skin and muscles showed numerous ecchymoses. The larynx and vocal cords very were œdematous, the air-passages were empty. The lungs retained their foetal condition, the colour being pale blue-red, there were numerous subpleural ecchymoses. The pericardium contained some fluid, and the heart was ecchymosed. (Nothing is said about the amount of blood contained by the heart.) The funis measured 66 cm. (26 inches) in length. The portion from the navel to the neck 15 cm. (6 inches), round the neck 15 cm. and from the neck to placenta 36 cm. (14 inches).

Hyperæmia of the brain and membranes is not a necessary accompaniment of death by strangulation, as the compression of the carotids impedes access of blood, indeed the condition of the brain and membranes is not constant, either in death from strangulation with the funis, or from compression of the funis occasioned by prolapse, out of twelve cases, Scanzoni found cerebral hyperæmia in four. The mark produced by strangulation with the umbilical cord is usually continuous round the neck its breadth corresponds to the thickness of the cord, the groove is rounded in transverse section and is soft, and what is very important, the skin is not excoriated.

Marks resembling those produced by the cord surrounding the neck may be caused by forcible bending of the head forwards during labour, especially in fat children, careful examination will show the absence of the signs of real compression, such as are produced by the application of a constricting medium. Forcible stretching of the neck may produce red stripes on the front or back of it. Kaltenbach<sup>1</sup> directs attention to these marks as resembling those due to strangulation, they disappear in two or three days with exfoliation of the cuticle. When the child's head has passed the os uteri, the pressure of the contracted cervix round its neck may leave a mark which will be broad and not so well defined as marks produced by other modes of constriction. The foetal membranes may become twisted round the child's neck and leave a suspicious-looking mark.

**Prolonged labour** is not an unfrequent cause of death to the child, the compression produced by a spastically contracted uterus after rupture of the membranes, if long continued, not only kills the child, but produces conditions of the body which may easily be mistaken for the results of intentional violence. Ashby,<sup>2</sup> in making post-mortem examinations of the bodies of recently-born infants, found meningeal hæmorrhage as a frequent result of asphyxia produced by prolonged labour. Stadtfelt<sup>3</sup> states that hæmorrhage in the cranium is liable to be regarded as a sign of criminal suffocation, whereas it is chiefly due

<sup>1</sup> *Centralbl. f. Gynecologie*, 1888

<sup>2</sup> *Brit. Med. Journ.*, 1890

<sup>3</sup> *Nord. Medicin. Arkiv*, Bd 17

to injury received in labour. The spine also may be injured by extreme bending, and blood found alongside it, in one case Stadtfelt found blood poured out about the kidneys without any obvious injury to the spine or any of the neighbouring organs. Effusion of blood is not uncommon when death of the child results from protracted labour, and this need occasion no surprise when the power of the uterine muscles and the delicate nature of the tissues of an infant at or before term are taken into consideration. In some cases the child lives several days after birth, although the injury which eventually caused death was then received. Keser<sup>1</sup> met with such a case in which he found eight ounces of blood in the peritoneum.

A remarkable case is recorded by Monteith<sup>2</sup> of a child, born after natural labour, with a depressed fracture in the middle of the right parietal bone, which was split from the sagittal suture on the one side of the depression to the coronal suture on the other, spiculæ projected from the inner surface of the fractured bone.

Deaths from injuries at birth increased from 448 in England and Wales in 1900, to 904 in 1911. In 1919 the number was 774.

**Immaturity or malnutrition** during intra-uterine life may lead to death of the infant shortly after birth, the latter is frequently due to placental degeneration, often of syphilitic origin, which interferes with the supply of blood to the fœtus, if the placental blood supply is very gradually diminished, as in progressive degeneration, there may be no signs of asphyxia in the fœtus, it is simply enfeebled from insufficient nutrition, if the circulation is more speedily interrupted by premature detachment of the placenta from the uterus, the signs of asphyxia will probably be present. The blood supply to the fœtus may be cut off by spontaneous rupture of the funis. Rivet<sup>3</sup> records a case in which a stream of blood from spontaneous rupture of the umbilical vessels followed breaking of the membranes.

Sudden death in new-born infants has resulted apparently from **hyperplasia of the thymus gland**. Grawitz<sup>4</sup> records two such cases, in one the gland covered the greatest part of the pericardium. Scheele<sup>5</sup> records another case in a child sixteen months old. The mode in which death is occasioned is not clear, although the post-mortem signs of asphyxia are present, it is possible, in some cases at least, that spasm of the larynx may have caused the asphyxia. Paltauf<sup>6</sup> states that there is no proof of compression of the trachea by hyperplastic enlargement of the thymus gland, which is simply one of the appearances due to general disease of the lymphatic system, and is not a direct cause of sudden death.

**Lethal malformation and disease**, as pneumonia or hydrothorax, are usually sufficiently obvious as to occasion no difficulty in arriving at a conclusion as to the cause of death. **Defective ossification** of the bones of the skull increases the risk of fatal compression during labour, and may be mistaken for injuries produced by violence. Taylor and Stevenson<sup>7</sup> mention the case of an infant whose body was discovered in a pond, presenting the appearance of injury to the skull in the form of two holes in one of the parietal bones, on careful examination it was found that the bone at the edges of the aperture was thinned down, and that originally the spaces had been covered with membrane which had disappeared with maceration.

<sup>1</sup> *The Lancet*, 1886

<sup>2</sup> *The Lancet*, 1874

<sup>3</sup> *Arch de Tocologie*, 1883

<sup>4</sup> *Deutsche med. Wochenschr.*, 1888

<sup>5</sup> *Zeitschr. f. klin. Med.*, 1890

<sup>6</sup> *Wiener klin. Wochenschr.*, 1890

<sup>7</sup> *Principles and Practice of Med. Jurisprudence*

Delayed parturition has been mentioned as a cause of mortality to the child, the opposite extreme—**hasty parturition**—is also attended with risk. The modes in which death may be thus caused are various. If the child is expelled head downwards on a hard surface the bones of the skull may be fractured. If the woman is standing upright when the child comes away the parietal bones may be broken, when but one is broken it is usually the left, on account of the rotation of the child in passing through the pelvis, the fractures may extend to the frontal or to the occipital bones. The child may be precipitated into a night commode or other utensil, and may be drowned in the liquor amnii or in other fluid into which it may fall. When the child falls into such a receptacle, it usually arrives head downwards, and, consequently, if not quickly removed a small quantity of fluid is sufficient to produce asphyxia. Rayner and Stuart<sup>1</sup> record a remarkable case. A primipara, aged 26, was left by her husband in good health and spirits at 2 p.m., and on his return at 4.30 was found dead in a water-closet. She had been delivered in this position, and the child's head had passed completely through the lower opening of the closet, the body and shoulders remaining above. Forcible attempts were made to extract the child, a healthy full-term female, but it was impossible to do so until the earthenware pan was smashed. An examination of the mother showed that the lower segment of the uterus and the cervix had been bilaterally ruptured through the whole thickness of the muscular wall, the mucous membrane and part of the muscular wall of the vagina had been fissured and the deeper tissues of the perineum lacerated. Superficially the perineum was intact. Death had been due to shock and hæmorrhage. The observers pointed out that from the medico-legal point of view the case is of interest, since had the mother not died it might have been argued that force other than the forces of expulsion and gravity had been used to get the child into the extraordinary position in which it was found. Mere falling to the floor without fracture of the skull may produce a fatal result from the injury sustained. Simple exposure to the cold air without covering and without any injury whatever, especially in the case of a weakly child, may be sufficient to cause death. In all these instances when lethal traumatic injuries are not produced by the fall, death results because the child receives no attention, if it is removed from the source of danger—drowning, or exposure to cold—it does not die. Usually the child is small or premature, and the head does not show a *caput succedaneum* or moulding.

In fatal cases the usual explanation why the child was left to die is that the mother became unconscious at the moment of delivery. This is a question of considerable importance, as it arises in a large number of cases of infanticide. Much scepticism is displayed by many experts as to the occurrence of unconsciousness during delivery. Heidenhain<sup>2</sup> relates the case of a woman seized with labour pains in the middle of the night, who got out of bed to obtain a light, when, as she alleged, the child was suddenly born, and as it fell to the ground the umbilical cord was torn, the mother returned to bed and fainted, and on regaining consciousness found that the child was dead, death being caused by intracranial hæmorrhage, the result of injury to the left parietal bone. Heidenhain looks with suspicion on the frequency of fainting after the birth of illegitimate children, and regards sudden labours in primiparæ as very unusual, both these objections have considerable weight, but cases beyond suspicion have occurred which prove the possibility of rapid delivery with simultaneous or subsequent insensibility. Pullmann<sup>3</sup> records an

<sup>1</sup> *The Lancet*, 1905.

<sup>2</sup> *Vierteljahrsschr. f. ger. Med.*, 1889.

<sup>3</sup> *Vierteljahrsschr. f. ger. Med.*, 1891.

instructive case of this kind. A married lady of good position, who anticipated with much pleasure the expected birth of a child, had some pains in the back, but did not regard them as labour pains. An hour or two after, bearing-down pains began, and the membranes ruptured, the doctor and nurse were immediately sent for. Meanwhile the patient felt a strong desire to urinate, and got out of bed to pass water, she placed herself on the vessel, but feeling a peculiar quick movement in the genital organs, she sprang up, and at the same moment the child fell into the vessel, the placenta following. The mother immediately lost consciousness, and if the nurse at that moment had not entered and removed the child from its perilous position head downwards in the vessel - it would undoubtedly have died. The child was in about the thirty-first week of development, there was no *caput succedaneum*. The mother had an extensive rupture of the perineum in consequence of the quick expulsion. If this had occurred to a servant-girl and in obedience to her desire to urinate she had gone to the closet, the child would have been dead long before she had recovered her senses, but her account of the event would probably have been regarded as a pure fabrication. Brunon<sup>1</sup> relates the case of a primipara near term who felt some pain in the back and a desire to defæcate, she essayed to do so and then returned to bed, on feeling a still more imperative desire she again raised herself in order to go to the commode, but was surprised to find something between her thighs, which was the head of the child. She knew nothing of what had taken place until she became aware of the presence of something externally. In another case reported by Langier,<sup>2</sup> a woman eight months advanced in pregnancy was seized with an irresistible desire to relieve the bowels and on getting out of bed the child was suddenly expelled into the utensil the funis was not torn, and the placenta had to be removed subsequently, the child was resuscitated. The woman felt nothing until a quarter of an hour before expulsion. Ehrendorfer<sup>3</sup> records a case of hasty labour that occurred in a lying-in hospital. A multipara in the eighth month of pregnancy for some time had complained of abdominal pains, she went one night to the closet, and in ten minutes returned stating that the child had come from her, the placenta fell from her as she arrived in the delivery-waid, the attached part of the cord measuring 10 inches, the free end being torn. The body of the child was found in the closet, it was 19 inches long and 4½ inches across the shoulders, a *caput succedaneum* was present, the lungs contained fluid like that in the closet. The woman died seven days after, and at the autopsy tuberculosis of the spine and a psoas abscess were found, which explained the constant pain experienced by the patient, and made her assertion that she did not know when the labour pains came on extremely probable. These cases show that statements made under circumstances of great suspicion are not, therefore, to be summarily rejected as being all but impossible.

When a woman becomes unconscious at the moment of expulsion, she must on regaining consciousness perceive what has taken place, and consequently imparts a guilty appearance to the event if she conceals the dead child and keeps the matter secret. The body of the child should be carefully examined for signs of injury inconsistent with the statements made, and the contents of the air-passages should be examined for fluid or other substance of the nature of that into which it is alleged the child fell. The funis, if torn, should be examined for indications of artificial separation, either with a cutting instrument or by being pulled asunder with the hands. If the infant was vigorous

<sup>1</sup> *Journ de Med et de Chirurg.*, 1890

<sup>2</sup> *Annales d'Hygiène*, 1891

<sup>3</sup> *Wiener klin Wochenschr.*, 1895

it might, in the act of falling into the receptacle, inspire sufficient air to make the lungs partially buoyant, in any case, indications of asphyxia would be present. In homicidal cases the child is usually suffocated or strangled, and consequently the body should be specially examined about the mouth, nostrils and throat.

**Hæmorrhage** from the funis, divided after being ligatured, may cause death, apparently from indispension of the blood to coagulate, or from insufficient tightness of the ligature.

### DEATH OF THE INFANT FROM CRIMINAL VIOLENCE.

The modes of death now to be considered are those which constitute the crime of infanticide. It is obvious that an infant is entirely at the mercy of any one disposed to take its life, there is scarcely any limit, therefore, to the ways by which this may be accomplished. Certain methods, however, preponderate in frequency, being easy of execution and because they most nearly simulate the appearances which are met with after death from accident or from natural causes.

**Suffocation.** In 1919, twelve deaths of infants under one year of age were attributed to murder by suffocation in England and Wales. A newly-born infant is easily suffocated by the application to the mouth and nostrils of any soft substance—as the bare hand, or a folded piece of cloth. Although very slight pressure is sufficient for the purpose, if it is applied directly to the mouth, the tender tissues of the infant usually yield evidence of its occurrence. It is very rare that the minimum amount of violence necessary to accomplish the object is used—it may be accepted as an axiom that in almost all homicidal attempts more violence is used than is sufficient to cause death. This applies with exceptional force to child-murder—the perpetrator is usually a woman who is under the influence of strong emotion at the time she commits the crime, and, therefore, is not in a condition to gauge nicely the degree of violence requisite to cause death, another reason why excessive violence is often used is the necessity of immediately silencing the cries of the infant. The external appearances after death vary with the mode in which suffocation was effected—whether by pressure directly applied to the mouth or by enveloping the infant in some soft material which prevents access of air. Fielitz<sup>1</sup> relates three cases of infanticide by suffocation. The first was accomplished by covering the child with a feather-bed, in this case no external signs of violence were present. In the second case the mother suffocated the child by placing her hand over its mouth and nose. At the inspection the mouth was found closed, the lips white, with the point of the tongue between them, four marks corresponding with the finger-nails were found on the right side of the neck just above the shoulders, and a single mark on the left side in a similar position, the internal signs of death from asphyxia were present. Immediately after the child was born the mother placed her left hand over its mouth and thus prevented expiration—the marks on the throat being caused by the finger and thumb nails resting on the neck, the fingers and thumb having been flexed so as to adapt the hand to the convexity of the child's face. The third case showed no trace of violence in the neighbourhood of mouth and neck, nor on any other part of the body, internally the signs of asphyxia were visible. Suffocation had been induced by enveloping the child in an apron. Schiller<sup>2</sup> made an examination of the body of a new-born infant, and found, in addition to the signs of

<sup>1</sup> *Vierteljahrsschr. f. ger. Med.*, 1891

<sup>2</sup> *Vierteljahrsschr. f. ger. Med.*, 1887

asphyxia, a quantity of blood in the finer bronchi. At the back of the pharynx he found injuries which had been inflicted by the mother having pushed her forefinger into the child's mouth with the object of producing suffocation, the finger had been firmly held there for some time, and the finger-nail had excoriated the pharyngeal wall and so produced the hæmorrhage. Suffocation was due to the presence of the finger rather than to the blood in the bronchi, as the blood was unmixed with air.

The body of an infant that has died from suffocation may be found in a medium capable of causing suffocation, death having been produced before immersion, hence the necessity of not accepting a probable cause of death without due investigation. The body may be found in a pond, or in a cesspool, but the child may have been dead when it was deposited there. Scrupulous care should be taken to search both the air-passages and the intestinal tract for traces of the medium in which the body was found, if traces are present they afford evidence of life at the time of immersion. A solid medium in a divided state as sand or loose earth may yield similar evidence of burial alive. Maschka<sup>1</sup> records a case in which the body of an infant was found buried about a foot deep in a field. The mouth contained a mass of mud-like consistence composed of sand and earth such as surrounded the body, some of which had penetrated as far as the pharynx and larynx, but could not be traced further neither into trachea nor œsophagus. The stomach contained no trace of it, but in the duodenum a small fragment was found. The lungs afforded proof of respiration, and presumptive indications of death from asphyxia were present. The opinion given was that the child breathed before being placed in the earth, and that by subsequent attempts at inspiration some of the soil was drawn into the mouth, but was too coherent to pass further, in the movements of swallowing which accompanied the attempts at breathing, a fragment of the soil found its way into the stomach, and was carried forward into the duodenum by peristaltic movements which do not immediately cease at the moment of somatic death. Tardieu<sup>2</sup> records the discovery of an infant's dead body in a box, the child having died from suffocation, but not from being shut up in the box, the under lip was everted and imprinted with the texture of a woven fabric, a fragment of which adhered to it, suffocation had been effected by pressure of a linen cloth on the child's mouth.

As previously stated, a new-born infant is easily suffocated, leaving it under the bed-clothes is sufficient to extinguish life without producing any of the external signs of suffocation. In such cases, due reserve must be exercised in attributing death to criminal intent, the mother may have fainted or otherwise have become unconscious, or irresponsible for her actions, for a period sufficiently long as to allow of the death of the infant without having entertained any design of taking its life. No hard-and-fast rule can be laid down. In married women who have just undergone the pangs of labour, the capacity for appreciating external conditions varies to an almost unlimited extent, some women become maniacal at the moment of delivery, and subsequently lapse into a semi-unconscious state, others are apathetic as to the fate of their offspring, for a time quite sufficient to allow of death from neglect if the requisite attention is not forthcoming, yet subsequently display the liveliest interest in its well-being. The opposite extreme, usually met with in unmarried women, is equally well marked, it is no uncommon event for a servant-girl to be delivered of a child in the same room with her fellow-servant, and to

<sup>1</sup> *Ibid*, 1886

<sup>2</sup> *Étude Médico légale sur l'Infanticide*, 1868



dispose of its body without exciting any immediate suspicion. In one case a servant-girl hearing the front-door bell ring at the moment the child was born instantly attended to the door and let her mistress in, it was subsequently found that at the time the placenta was still within the uterus, and that hæmorrhage was taking place. In another case a servant-girl went a distance from home, and when returning she asked the boy who drove the cart in which she rode to stop, she got out, went to a recess in the hedge, and five minutes after was seen walking home, a distance of a mile and a half, the day following she went about her work as usual. In the five minutes' interval she had given birth to a living child. The external influences which affect married and unmarried women respectively, as to their behaviour during and after labour, are to be taken into consideration in forming an opinion as to the probability or otherwise of intentional neglect on the part of the mother of anything necessary for the well-being of the infant. As regards unmarried women, there are *prima facie* grounds for suspecting them to be desirous of the death of the infant to which they have given birth, but in the absence of incriminating circumstances it is unjust to turn a deaf ear to the woman's statement— that she lost consciousness at the moment of delivery. Since it is a fact that married women who desire their children to live not infrequently succumb for a time to the violence of their emotions, or of their sufferings, and imperil the life of their offspring, the possibility of a like condition attacking an unmarried woman is not to be denied.

**Strangulation.** In England and Wales, the deaths of 24 children under one year were officially stated to have been due to murder by strangulation in 1919. The number of deaths from this cause has fallen substantially during the last decade, the annual average from 1905 to 1911 having been 86.

Strangulation may be effected either by compression of the child's throat with the unaided hands, or by means of a cord, ribbon, or other similar medium. When strangulation is effected with the **unaided hand**, marks produced by the pressure of the fingers are usually to be seen, it frequently happens that two or more impressions produced by the finger-nails are found on the left side of the child's neck, and a single impression on the right side. This arises from the throat being clutched between the fingers and thumb of the right hand. The nail-marks may be further back than might be expected, on account of the whole neck being grasped instead of the air-passage only. In addition to the marks of the nails there may be superficial ecchymoses more or less corresponding to the lines of pressure produced by the fingers, the ecchymoses often spread to the ears and along the sides of the face, excoriations of the cuticle produced by scratches of the finger-nails may be present. In some instances no external signs are visible, although death has been produced by throttling, in these cases it is probable that the web between the thumb and first finger was applied to the front of the throat, the back of the neck being supported by the palm of the other hand, the intervention of a soft fabric between the hand and throat materially lessens the amount of external marking. Although no superficial ecchymoses are visible, the deep structures may be infiltrated with blood. If great violence has been used the muscles are bruised, the cartilages of the larynx probably fractured, and even the vertebræ may be injured.

Marks resembling those produced by criminal violence may result from attempts on the part of the mother to aid her delivery. Scratches produced by the finger-nails may be found in the neighbourhood of the nose, mouth, and ears, with possibly indications of pressure by the finger-tips on the under

part of the lower jaw and side of the neck. It is exceedingly unlikely that the mother could endanger the child's life by attempts to expedite delivery, when severe injuries have been inflicted, the presumption is against this explanation. In arm or leg presentation, fracture or dislocation might be caused by improper traction on the part of the mother in her attempt to drag out the child.

Suffocation may be produced by firm, prolonged compression of the chest and abdomen so as to prevent inspiration, unless a soft fabric protected the child's body, the marks of the finger and finger-nails would probably be distinguishable. Tardieu mentions such a case in which there was a depression in the chest and abdomen.

When a piece of cord or ribbon, or other similar medium, has been used to strangle the child, a more or less distinct groove will be found round, or partly round, the neck. The breadth and depth of the groove, within limits, is governed by the kind of material used, and the degree of force applied, a thin cord applied with the same degree of force would produce a narrower, deeper furrow than a handkerchief. The bottom of the groove is usually white or grey, the borders being livid or of a violet colour. There will probably be some superficial ecchymoses visible in parts of the groove or its borders, but not invariably so. Excoriations of the epidermis are likely to occur, unless the strangulating medium was very smooth and soft in texture. Deep extravasations of blood below the groove and around it are almost invariable, and if great violence has been used there may be laceration of the muscles, with injuries to the larynx or trachea, and even to the cervical vertebræ. Internal examination reveals the signs of death from asphyxia.

The usual defence in accusations of infanticide by strangulation is—that death was caused by the funis surrounding the neck during labour. The indications of accidental strangulation produced in this way have already been described, one or two additional points remain for discussion. First, as to the appearance of the mark round the neck. It has been erroneously assumed that local ecchymoses are not present when the funis is the constricting medium, but the results of post-mortem examinations in undoubted cases of accidental strangulation with the funis prove the contrary, there is, however, a distinction (previously mentioned) which is of considerable moment as a diagnostic sign—the smooth, soft funis does not excoriate the skin. Excoriation of the skin is not invariable in homicidal strangulation, but when present it points in that direction. When the funis encircles the child's neck at birth it not unfrequently goes more than once round, which will be indicated by the mark on the neck. As a rule, in cases of accidental strangulation caused by the coiling of the funis round the child's neck, the lungs will be found in the foetal state—they will sink when placed in water. This rule is subject to exceptions. The asphyxia which causes death in these cases may result from arrest of circulation in the umbilical vessels as well as from constriction of the child's neck. When the head is expelled the constriction may be just insufficient to cause strangulation, although the tension on the funis may be sufficient to arrest the circulation through it, which would determine the commencement of respiration, very little additional tension would arrest the breathing, and in the absence of skilled assistance the child would die during birth. On examination the neck would show signs of strangulation, and the lungs would yield evidence of having respired. Winter<sup>1</sup> records a case illustrating this complication. A woman, aged twenty-one, with normal pelvis, was attended in labour by a midwife,

<sup>1</sup> *Bericht über d. Sitz. d. Ges. f. Geb. u. Gyn.*, 1885

the funis surrounded the child's neck too tightly to be removed, and the head was expelled, but the shoulders impeded further progress, a medical man arrived in five minutes and he tied and divided the funis. The child was delivered with considerable difficulty in a deeply asphyxiated condition, and only with prolonged efforts was it brought to life, a distinct strangulation-mark ran round the neck. Eight hours after, the child died. At the necropsy a clot of blood was found under the pia mater flattening the cerebral convolutions, blood was also found in the lateral ventricles and at the base of the brain. Ecchymoses were present on the pleura and pericardium, and extravasation of blood under the mucous membrane of the larynx, the lungs were partially distended with air. In the absence of any history it would probably have been assumed that the child was born alive and was afterwards strangled.

The average length of the funis is about 20 inches. It may not exceed 6 inches, or it may reach 70 inches or more. When both foetal and placental portions of the funis are forthcoming they should be measured, the total length might be insufficient to allow of the child's neck being encircled by it, further, the tension-limit of the funis should be tested, it might be unable to bear the strain necessary to strangle the child. The funis has been used as a medium for **criminal strangulation**, the allegation made being that it surrounded the child's neck at birth. In such cases the cord will be stretched, and will probably show traces of local displacement of the Wharton's jelly from being twisted round the fingers in the act of strangulation, the funis is so smooth and slippery that simply grasping it with the hand scarcely affords the necessary hold. The lungs would probably show signs of having partially respired, although in one case of criminal strangulation with the funis the lungs were found in their foetal state.

Strangulation-marks round the child's neck are sometimes alleged to have been produced by a handkerchief or piece of tape with which the mother attempted to aid her delivery, this is an improbable supposition, and in any case very little force could be thus applied, so that careful examination of the marks externally, and dissection of the deeper structures underlying them, would probably enable a conclusion to be arrived at.

Intentional **neglect in tying the funis** may lead to death from hæmorrhage. The absence of a ligature does not prove that the cord was not tied, it may have been removed, or the funis may have separated at the site of the ligature. This warning is specially appropriate in the case of infants whose bodies are not examined until a week or two after death, post-mortem softening may easily cause detachment of the distal end of the funis along with the ligature. Devergie<sup>1</sup> relates a case in which an infant with cord and placenta was being removed from a river where it had remained for about a fortnight, the funis was so tender that it gave way, and the placenta was carried down the stream, a medical man, not knowing these facts, examined the body, and concluded that the funis had been torn at the birth, and that death had resulted from hæmorrhage. The free end of the funis should be examined as to the condition of the structures which enter into its composition. If it has been divided with a sharp cutting instrument, the transverse section will be clean and even, if it has been torn the section will be jagged and irregular, some of the structures being more elastic and less yielding than others, if division had been effected by means of a blunt knife, it will probably be difficult to determine the mode of separation, in which case it is well to avoid giving a decided opinion.

The importance of the question as to how the funis was divided is well

<sup>1</sup> *Annales d'Hygiène*, 1873.

illustrated by the following case narrated by Koch<sup>1</sup>—A servant-girl, advanced in pregnancy, felt a desire to relieve the bowels, she went to the closet and was suddenly delivered, the child falling through the opening, the girl explained the event as being accidental. The placenta subsequently came away, and the child was removed from the closet asphyxiated. The funis was nearly 20 inches long—it was of moderate thickness, and was separated about the mid portion of its length—10 inches from the navel. On the foetal portion there were three injuries, two of which affected the amniotic sheath only, the arteries not being injured—the third went half through the funis, one artery and the vein being divided, the injuries were comparatively smooth, except the one through the amniotic sheath, which was rather ragged, the vessels were irregularly divided, and they partially projected beyond the other structures. Two experts pronounced that the funis had been divided with a sharp cutting instrument, basing their opinion on the presence of the injuries described, and on the unusual position of the point of separation, accidental rupture usually taking place either close to the placenta or close to the navel. Re-examination led to a change of opinion—that the injuries were not produced by a cutting instrument, but by the sudden tug of the falling child. Winckel states that he has often seen injuries of this kind produced by sudden tension of the funis. The medical expert witness should ever be alive to the importance of not accepting what appears at first sight to be an evident explanation of the cause of injury, without considering whether there may not be another and correct though less obvious interpretation. In the above case the first opinion, if accepted, would have condemned the girl as a murderer, the second to some extent confirmed her story, and established the possibility of the affair being really accidental. According to Winckel,<sup>2</sup> rupture of the funis from precipitate labour, in more than three-fourths of the cases, takes place within 6 inches from the umbilicus, in 12 per cent, close to or within the umbilicus—still less frequency in the middle of the cord or near the placenta.

**Fractures of the skull** are frequently found to have been inflicted on newborn infants when death is suspected to be due to infanticide. When the child is killed by blows on the head, or by dashing it against a wall, the injuries inflicted are usually excessive. It is a mode of killing which does not admit of the attempt to attribute death to natural, and scarcely to accidental, causes, the sole object is to deprive the child of life as quickly as possible, and to trust to concealment of its body in order to escape the consequences, the result almost invariably is that the bones of the skull are literally smashed in. When the deed is perpetrated by the mother she is usually at the time in a state of maniacal excitement, and is, therefore, in no condition to pay heed to the amount of damage she inflicts. After the child is dead it is generally thrown into a pond or cesspool, or is buried in the ground.

The defence usually is that the injuries were caused at birth from difficult or from precipitate labour. As previously shown, fractures of the bones of the skull may be caused during ordinary difficult labour—without the use of instruments or of manual interference—and, further, they may present an appearance not unlike that which results from modified criminal violence. When the injuries have been intentionally inflicted there will be more or less damage done to the external soft parts, the scalp will show signs of the force resorted to, in the form of excoriations, rents, or complete disintegration of structure, which are not seen when the injury to the skull has been caused during ordinary non-instrumental labour. If fracture has resulted from the head being forced through

<sup>1</sup> *Arch. f. Gynæcologie*, 1886.

<sup>2</sup> *Lehrbuch d. Geburtshilfe*, 1889.

a contracted pelvis, the shape of the head and the presence of a well-developed cephalhæmatoma will afford evidence of the fact. The presence of blood under the scalp or the pericranium, or even within the skull, is quite consistent with natural causation. When criminal violence has been resorted to, there will probably be slight injuries visible on other parts of the body besides the head, such as finger-marks, scratches, and bruises, from rough handling during commission of the crime. If, on the other hand, the defence is that the woman was **suddenly delivered** whilst in the **erect posture**, attention should be directed to the point of separation of the funis, usually, but not invariably, it is near the navel, in order to determine whether it was cut or torn asunder, the free end of the funis is to be carefully examined with a lens. The injuries sustained by the skull will probably be much less than those criminally produced. Whilst it is undoubtedly true that the child's skull may be fractured by the mother being delivered in the upright posture, the child falling head downwards on to a hard surface, yet hasty delivery has frequently taken place under these conditions without the child receiving material damage. Snell<sup>1</sup> records the case of a woman who in the upright posture was unexpectedly delivered of a child at term, the child fell head downwards on a stone floor, rupturing the funis in its fall, yet it sustained no injury, not even a bruise being apparent. If the injuries to the skull are extensive, homicide is indicated, lesser degrees of violence are compatible with either criminal or accidental causation, and consequently demand careful consideration of all available indications, and frequently no slight discrimination in order to interpret them aright.

Another defence may be made—that the injuries to the skull were accidentally inflicted by the mother in attempting to expedite her delivery. It would be difficult to fracture the skull in this way, though it would not be impossible. The degree of injury inflicted on the skull, together with the absence of marks of violence on the rest of the body, are the indications from which an opinion is to be formed.

Sometimes the dead body of an infant is found on a waste piece of ground surrounded by a wall, and it becomes a question as to whether any injuries sustained by the head may not have been caused after death by the body being thrown over the wall, the same consideration is involved if the body is found under a heap of stone, or in a coal-heap, or if it has been forced down a soil-pipe or drain. The great difficulty in answering this question arises from the fact that an injury caused *immediately* after death is not characteristically different from one inflicted during life. It was formerly assumed that a blood-stained appearance of the edges of the fractured bones indicated that the injuries had been inflicted during life, but if the injuries are produced soon after death a like appearance ensues, if some hours elapse between death and the production of the fractures the absence of such signs is to be expected. No positive opinion can be formed if the edges of the fractured bones are blood-stained, as to whether the fractures were made before or immediately after death, if there is no staining of the edges, and but little effusion of blood within or without the skull, the inference to be drawn is that the injuries were caused after death.

Violence inflicted on the child's neck, producing subluxation, or fracture of the cervical vertebræ, may have caused death, evidence of the injury will be observable on examination of the body.

**Wounds** produced by cutting or pointed instruments are frequently met with on the bodies of infants found dead. They vary in extent from minute

<sup>1</sup> *Brit Med Journ*, 1891

punctures penetrating vital organs, to extensive wounds such as result from decapitation, or even complete dismemberment. Punctures through the fontanelles, the orbital plate, between the vertebræ, and between the ribs into the heart, have been discovered after death, these less apparent methods of causing death should be borne in mind when no obvious injury can be discovered. The more formidable injuries reveal themselves at first sight, the difficulty in dealing with them arises from the usual defence—that they were inflicted after death, with the object of facilitating the disposal of the body. The signs by which a diagnosis is to be made as to whether wounds have been inflicted before or after death are elsewhere described. In the case of smaller wounds it may be asked whether they were sufficient to cause death. Next to injury of a vital organ, hæmorrhage has to be considered as a cause of death, an exsanguineous condition of the body and internal organs generally, points to hæmorrhage. The condition of the funis—whether it has been tied or not—should be examined before an opinion is formed as to the probability of death having occurred through loss of blood from wounds. If it is alleged that the child was still-born, and that it was mutilated with the object of more easily concealing its body, the presence or absence of the signs of respiration, as far as they are available, must be ascertained.

An instance of mutilation of the child during birth is given by Barbour.<sup>1</sup> An unmarried woman travelling alone in a railway carriage was suddenly seized with labour pains, feeling something protruding from the vagina, she attempted with some force to deliver herself, and in so doing broke the presenting limb—an arm—which in a moment of pain and frenzy she then severed with a table knife and threw out of the window. On reaching home she was delivered by version of a still-born well-nourished male child, with the right arm nussing two inches above the elbow, a considerable quantity of blood had escaped from the vagina after the amputation of the arm, and the child's body was palid, so that hæmorrhage was doubtless the cause of death.

**Drowning** was the method of murdering 17 infants under one year of age in England and Wales, in 1919. When a body is found in water it must be remembered that the bodies of infants already dead are often thrown into ponds, canals or rivers to get rid of them. An infant's body, if well nourished, contains a large proportion of fat, and consequently does not readily sink. If the body is found in water with a mummified funis it is certain that the child, living or dead, remained in air sufficiently long for mummification to take place—a funis once mummified does not return to its original condition on being placed in water. On the other hand, if the child is submerged, either living or dead, while the funis is fresh and plump, mummification does not occur. The signs of death from drowning do not differ in the infant from those found in the adult, they will be described elsewhere. If still-birth is alleged with the object of reducing the crime to concealment of birth, an examination of the lungs will determine the question unless putrefaction prevents, in the absence of clear evidence of death from drowning careful search should be made for other causes of death. When the body is found in a night commode or closet, and hasty parturition is tendered as an explanation, the funis must be carefully examined with the object of ascertaining whether it has been torn asunder or divided with a cutting instrument. It has been stated that in the act of falling the child may make an inspiration sufficiently vigorous as to fully inflate the lungs, but from what has been already said on this subject, it may be inferred that complete inflation could not take place under such circumstances, it is

<sup>1</sup> *The Lancet*, 1892

not to be denied, however, that partial inflation might occur. Any fluid or other matter found in the air-passages or stomach should be compared with the medium in which it is alleged the child was drowned.

A superficial glance at a section of the lung might lead to the assumption that no fluid was present, as it is not always found in the larger bronchi. There is an absence of frothy mucus in the lungs of new-born children that have been plunged into a liquid before breathing, under these conditions an infant makes attempts at respiration and inspires some of the liquid, but as no air is already in the lungs, and none enters with the liquid, the bronchi do not contain froth such as is met with after drowning in the lungs of those who have respired. Dittrich<sup>1</sup> records the case of a pregnant woman who suddenly felt a desire to defæcate, and in consequence placed herself over a small tub containing waste water, coffee-grounds, dust, and the like, she immediately gave birth to an infant at or near term, and tore the funis without tying it. On section of the child's body the lungs were found in the fœtal state, except that they contained much blood, they sank both whole and divided. In the trachea and larger bronchi no foreign substance was visible, but on careful examination the lumina of the small bronchi were seen to be filled with a mixture which corresponded with the medium in which the infant was drowned, on microscopical examination the finest bronchioles were found to be choked up with it. The stomach and intestines sank when placed in water.

Intra-uterine maceration must not be lost sight of as a possible cause of changes resembling those produced by prolonged submersion in water.

Destruction of the body of a new-born infant is sometimes attempted by **incremation**. The question in such cases are—Was the child still-born? If not, was it alive or dead when subjected to the action of fire? The first question is to be answered in accordance with the signs already given, and the second with those described in the chapter on death by burning.

An advanced stage of incremation may make it impossible to answer either question; the problem then becomes—Are the remains those of a human fœtus or not? One of the bones may possibly afford the necessary evidence, as in the following case—A charred, shapeless mass was found on a kitchen fire, when withdrawn it measured  $9\frac{1}{2}$  inches in length and 6 inches in its greatest diameter, it weighed 4 ounces. On examination indications of ribs, spinal column (much twisted) and base of the skull were found, but nothing to distinguish the remains from those of an inferior animal until, on lifting up a projecting fragment of carbon, an intact scapula was discovered, this proved that the remains were human, as the postero spinal segment of the bone was much longer than the anterior, the difference in the lower animals being either insignificant or entirely absent. A servant girl who was suspected of having been recently delivered subsequently confessed that the remains were those of a fœtus at about the fifth month.

**Exposure to cold and deprivation of food**, if sufficiently prolonged, will, of course, cause death of the infant. Deliberate infanticide is not often attempted in this way, when a child is exposed so as to endanger its life it is with the object of getting rid of it without actively committing murder. The defence usually offered is— that the child was born unexpectedly (hasty parturition), and that it was still-born, or if living, that it was exposed so as to attract attention (before the door of a house, for example) in order that it might be rescued. In like manner a new-born living infant has been placed in a hamper with a feeding-bottle and sent by train to the putative father.

**Poisoning** is rarely resorted to in the newly born as a means of infanticide, if suspected, an analysis of the digestive organs and their contents must be made.

<sup>1</sup> *Prager med. Wochenschr.*, 1890.

### 5. THE LENGTH OF TIME THE CHILD HAS BEEN DEAD.

This is to be estimated by the stage at which the post-mortem phenomena have arrived. The tendency is for animal heat and cadaveric rigidity to pass away, and for putrefaction to occur sooner than is the case with the adult body. When the putrefactive stage is reached no positive opinion is justifiable as to how long the child has been dead.

### POST-MORTEM EXAMINATIONS IN CASES OF SUSPECTED INFANTICIDE.

When making a post-mortem examination of the body of an infant suspected to have been murdered, it is necessary to investigate very closely the **external appearances**. The presence of body-warmth, cadaveric rigidity, or of the signs of putrefaction should first be noted. If the body has been found without any history, a preliminary search should be made for marks or other indications which might lead to identification, and notes should be made of articles of clothing, or of the box, paper, or other envelope within which the body was found. *Vernix caseosa* is to be sought for, its presence demonstrating absence of the usual attentions at birth. The funis should be carefully examined as to the mode in which it has been separated, the portion attached to the child's abdomen should be measured. The whole of the surface of the body must be examined for scratches, bruises, and other slighter injuries, if any are found, attention should be directed to their probable causation—from accidental causes during or after birth, or from criminal violence. Before touching the mouth the position of the tongue and lips should be observed—whether the tongue protrudes, and the lips are flattened or everted. Any other indications of pressure on the mouth and nostrils—such as might be produced by the bare hand, or by the application of a cloth—with the presence of punctiform ecchymoses on the face, and its general hue, whether pallid or livid, should be noted. The natural apertures of the body should be examined for foreign bodies. Any marks on or round the neck must be closely investigated, and the underlying structures subsequently examined by dissection. Should the whole body be extremely pallid, observe if there is any external cause for hæmorrhage other than an untied funis. The length of the child, its weight and state of nutrition should be noted, bearing in mind that the dead body of a new-born child tends rapidly to lose weight by cutaneous transpiration, especially when exposed without covering, or merely enveloped in paper or placed in a box.

**The internal examination** is commenced by making the usual incision through the skin from the upper part of the manubrium to the pubes. After opening the abdominal cavity, the forefinger of the right hand is introduced and its tip placed on the highest part of the arch of the diaphragm, with the forefinger of the left hand the intercostal spaces are counted from above downwards until the position of the two fingers corresponds—this determines the height of the diaphragm. The thorax is then opened and the condition of the heart ascertained *in situ*. It is convenient to continue the primary incision upwards through the skin of the neck and the lower jaw, and to divide the symphysis with a sharp-pointed pair of scissors, passing one blade within the mouth, from above downwards in the middle line, immediately behind the alveolar process. Separate the soft structures and fold the two halves of the jaw outwards and backwards, in this way a complete view of the mouth and pharynx is obtained without disturbing any foreign body that may be present, the



tongue can be drawn forwards or to one side, as is most convenient. The lungs, stomach, and intestines are removed for testing as to specific gravity after their volume and general appearance have been noted. Before cutting through the scalp see if there is any trace of puncture, and after reflecting it examine the fontanelles with the same object, the possibility of puncture or other injury to the cervical cord should be borne in mind. The presence of meconium or of food-stuffs in the intestines is to be noted. The stage of development of the child must be estimated by an appeal to the various indications described in the section which deals with the subject. The existence and size of the ossific centre in the inferior epiphysis of the femur is ascertained by slicing away thin sections from the lower end of the bone until the maximum diameter of the centre is reached. The possibility of poisoning must not be lost sight of, the organs necessary for chemical analysis should be removed for that purpose if deemed necessary.

**Concealment of Birth**—"If any woman shall be delivered of a child, every person who shall by any secret disposition of the dead body of the said child, whether such child died before, at, or after its birth, endeavour to conceal the birth thereof, shall be guilty of a misdemeanour" (24 and 25 Vict, c 100, sec 60)

In a great many accusations of infanticide the jury reduces the crime to that of concealment of birth. Concealment of a living child that does not die before it is discovered, does not constitute the misdemeanour. In cases of concealment of birth the question of live birth does not arise, therefore medical evidence is limited to the following points - Are the remains those of a viable child? Has the accused been recently delivered? There is some uncertainty as to whether the secret disposal of a fœtus which has not reached a stage of development, compatible with viability, constitutes concealment of birth.

Stephen (*Crim. Law of Eng. 1891*) says that "the expression 'delivered of a child' does not include delivery of a fœtus which has not reached the period at which it might have been born alive." Erle J. directed the jury on a charge of concealment of birth (*R v Berriman*, 1854, 6 Cox, p 388) "that it must appear that the child had gone such a time in its mother's womb that it would, in the ordinary course of things, when born, have had a fair chance of life." Smith J. (*R v Hewitt*, 1866, 4 F & F p 1101) left it to the jury to say, "whether the offspring had so far matured as to become a child, or was only a fœtus or the unformed subject of a premature miscarriage." At the Chester Summer Assize, 1895 (*R v Walle*), Lawrence J. ruled in accordance with the opinion of Stephen above cited. On the other hand, Martin B. (*R v Colmer*, 1864, 9 Cox, p 506) said "that he saw nothing to limit the word 'child' in the statute to a child likely to live or likely to die, but as soon as the fœtus had the outward appearance of a child it was sufficient."

According to **Scotch law, concealment of pregnancy** is a crime. It is not necessary that the dead body of the child should be found, nor need there be proof of infanticide, all that the law requires is that the woman must be proved to have been pregnant sufficiently long as to render possible the birth of a living child. If, during her pregnancy, the woman reveals the fact to another person, even if it is with the object of arranging for concealment of the child's birth, and evidence to this effect is tendered, the law is evaded—the pregnancy is no longer "concealed."

## CHAPTER XVI

## BIRTH IN RELATION TO THE CIVIL LAW.

**Live birth**, in the legal acceptance of the term, is determined by the same conditions in both criminal and civil cases. In **criminal** cases the character of the medical evidence differs from that which is usually tendered in civil cases, because in the majority of criminal cases the woman gives birth to the child in secrecy, so that no witness to the fact that the child was fully born at the time when signs of life were developed is forthcoming, consequently expert evidence founded entirely on the indications presented by the dead body of the newly-born infant has to be relied on. As a matter of fact, unless there is distinct evidence of respiration having occurred, a charge of infanticide is practically never pressed. In **civil** cases witnesses are usually to hand—the accoucheur and the nurse—who were present when the child was born, and who therefore can testify to the fact of legal birth at the time the child displayed tokens of vitality, hence proofs of live birth are available in civil cases which, by circumstances, are excluded in criminal cases. The occurrence of respiration, to which so much significance is attached in cases of infanticide, constitutes but one indication of life, its importance arises from the fact that it leaves after death more or less permanent and reliable traces of its occurrence. With the exception of the stomach-bowel test, all other manifestations of a life that ceases almost immediately after it has demonstrated its existence, vanish and leave no trace behind.

There is no statutory definition of live birth, and the law on the subject is accordingly derived from judicial decisions. In England, the most evanescent sign of life, **provided that** it is observed after the **child is entirely outside the body of the mother**, constitutes proof of live birth. With this proviso the least muscular movement that may amount to a mere momentary twitching of the lips or of a limb, or a few beats of the heart, either heard with the stethoscope or felt in the cardiac region, or by pulsation in the undivided funis, or the more obvious evidence afforded by respiration, with or without crying, are each, and severally, proofs of live birth. The duration of any of these signs is immaterial—one moment of life thus manifested avails, as proof of live birth, as much as though the child continued to live for days or months. According to the law of Scotland the commencement of respiration after expulsion must be proved to establish live birth.

In civil law, live birth comes into consideration in respect to the inheritance of property, in such cases much may depend on the fact that the child displayed, or did not display signs of life after birth, and in the event of live birth, on proof of the exact time when it was born. If the head is born, and an interval elapses before the body is expelled, during which the child is observed to breathe and cry, that is not the moment of birth. The delay may extend over some minutes, and, as the law does not recognise the fractional part of a day, if it occurs partly before and partly after midnight, a mistake in recording the exact time of birth might make a difference of twenty-four hours, and possibly

alter the succession to an estate if the child continued to breathe until and after birth. It may die, however, during the delay, before the body is expelled, in which case the child is still-born, the law taking no cognisance of signs of life manifested before complete expulsion.

The question of live birth has to be considered in respect to **Tenancy by Courtesy**. "When a man marries a woman seized of an estate of inheritance, and has by her issue born alive, which was capable of inheriting her estate, in this case he shall, on the death of his wife, hold the lands for his life as tenant by the courtesy of England."

It will be seen that there are two obvious conditions to be fulfilled—(a) That the issue must be borne alive, (b) that it was capable of inheriting. A third condition, which is not quite so obvious on glancing over the above quotation, is (c) that the woman must be living at the time she gives birth to the child.

(a) The proof of live birth is afforded by the evidence of the medical man, or of the nurse, or of both, who were in attendance at the labour. (b) Capacity for inheritance requires that the child shall have the shape of mankind. This question has been previously discussed under the head of sexual and other abnormalities. (c) Death of either wife or husband dissolves their marriage. If a married woman who is in possession of an estate held on the above-named terms is pregnant with a living child and dies before the child is born, her estate at once descends to the heir-at-law, not being intercepted, as it were, by the birth of the child. If the birth of the child precedes the death of the mother by ever so little, the child being born alive but dying immediately, the husband acquires the estate for the rest of his life. By Cæsarean section a child may be removed alive from the body of its *dead* mother, in such a case the husband would *not* become tenant. The point has been discussed as to whether a child extracted by Cæsarean section from the body of a *living* woman can be regarded in the legal sense as being born, it is probable that under these conditions the husband could enjoy the tenancy.

**Notification of Births.**—Under the Notification of Births Act, 1907, the birth of a child which has issued forth from its mother after the expiration of the twenty-eighth week of pregnancy must be notified to the medical officer of health of the district. Since 1915 this procedure has been obligatory in all districts. (For duties of medical practitioners in regard to this Act, see Chap XXV.)

**Still-birth.**—A still-birth need not be registered, but must be notified to the medical officer of health if of twenty-eight weeks' gestation. If it is desired to inter the child in a public burial-ground the authorities of the burial-ground must receive a written declaration of the mother's name, and of the fact of still-birth signed by a coroner or medical man who has seen the body, or by the person bringing it for burial. Usually the statement is given by the midwife or doctor to the undertaker, who then disposes of the body—sometimes, it is known, by placing it in the coffin of another person, a fact to be remembered in exhumations. This lax system obviously lends itself to the concealment of crime, or at least to the easy disposal of children who die shortly after birth.

## LEGITIMACY.

The law assumes every child born during marriage, or, after the death of the husband, within limits consistent with the normal duration of gestation, to be legitimate unless the contrary is proved.

**Proof** that the husband is **not** the father of the child may be adduced on the ground that he is **physically incompetent**, or that he had **not had access** to his wife within a period consistent with the date of the child's birth. Physical incompetency resulting from the extremes of age, or from malformation or disease, has already been dealt with. The plea of non-access, in addition to evidence as to fact—with which the medical witness has nothing to do—involves the consideration of the duration of (a) *normal gestation*, together with the questions how far may gestation be (b) *prolonged*, and how far may it be (c) *shortened* consistent with the birth of a living child.

(a) **Normal Duration of Gestation.**—There is no means of absolutely determining the duration of gestation, hence the discrepancies in the time allotted by different authorities. Of the many data from which the interval between impregnation of an ovum and the birth of an infant at term may be calculated, two only need be mentioned—the *cessation of menstruation*, and a *single coitus*.

The **cessation of menstruation**, apart from accidental interruption of the function, is an uncertain starting point. It is generally accepted that menstruation is not necessarily associated with ovulation—it may take place without ovulation, and ovulation may occur without menstruation. The impregnated ovum may embed itself in the uterine mucous membrane during any part of the intermenstrual period, and therefore the normal duration of pregnancy cannot be accurately determined from cessation of the menses. Matthews Duncan, reckoning from the last day of menstruation, computes the duration of gestation at 278 days, other authorities give a lower or a slightly higher computation.

A **single coitus** affords no fixed datum, because impregnation is not necessarily synchronous with it. Spermatozoa retain their activity within the female organs for many days, during any of which they may reach the ovule and fecundate it. Computed from a single coitus, the duration of gestation is stated by Ahlfeld<sup>1</sup> to be 271 days, by Stadfeldt<sup>2</sup> 272 days, by Duncan<sup>3</sup> 275 days, according to Lowenhardt<sup>4</sup> the average from statistics gives 272 days.

The **normal duration of pregnancy** is generally accepted as being about **280 days**, with the possibility of exceeding this period. The duration of pregnancy should always be expressed in days, months may be either calendar or lunar, therefore the use of the word is conducive to error.

(b) **Abnormally Prolonged Gestation.**—Attempts to estimate the extreme limits of gestation in either direction are further embarrassed by the fact that, in almost all such cases, the evidence on which the estimations are based is liable to be tainted. In many cases the interests of the individuals concerned are profoundly involved either from the moral or the pecuniary standpoint, and apart from wilful misrepresentation, women are very apt to be misled by their feelings and wishes. It may be premised that the usually accepted duration of gestation is undoubtedly capable of considerable extension, the difficulty lies in limiting the degree of extension. In some countries the limit is fixed by law—in France and Italy the limit is 300 days, in Germany a child born within 302 days after the death of the husband is regarded as legitimate, in Scotland a child born within ten lunar months after the death of the husband is considered legitimate. In England and America no limit is fixed, and consequently the opinion of experts is taken, and the entire question is discussed at every trial into which the subject enters. The longest gestation yet allowed by the English courts is 307 days, in 1917 the court dismissed the petition

<sup>1</sup> *Monatsschr. f. Geburtshilfe*, 1869

<sup>2</sup> *Annales de Gynecol.*, 1877

<sup>3</sup> *Fecundity, Fertility, Sterility*, 1866

<sup>4</sup> *Arch. f. Gynakol.*, 1872

for a divorce of a husband who had been absent from his wife in another country for that period of time. The labour was normal, and the child of average size and weight. The petitioner was unable to support his case by other evidence of a conclusive character. In America, 317 days have been allowed in an indictment for seduction, and probably sentiment had something to do with the verdict, a claimant to an estate might have fared differently.

Many cases are recorded of gestation extending to 300 days and beyond, cases which approach, but do not exceed, 300 days have a much greater air of probability about them than those which claim additional weeks. Acker<sup>1</sup> reports a case in which gestation lasted 305 days after a single coitus, the child not exceeding in development that of full term, allowing for the uncertainty of the moment of impregnation after insemination, the duration of gestation in this case might be reduced to 290 days. In a case recorded by Purkhauer,<sup>2</sup> the woman menstruated in the non pregnant state regularly every 28 days. She had her last period on the 28th of April, 1889, she first felt the child about the middle of September, which would make the end of the first week in February, 1890, the probable date of labour. On the 13th of March, 1890, she was delivered of a living boy weighing 8 lbs 12½ ozs, and measuring nearly 21 inches in length. If conception is dated at seven days after the cessation of the last menstruation, the duration of gestation was 316 days, if seven days before the next anticipated period is taken—which, however, would not agree with the date of quickening—the duration of gestation would be 300 days. Thomson<sup>3</sup> records a case in which the duration of gestation, reckoned from the last day of menstruation, was 317 days, and from the last coitus 301 days. Duncan<sup>4</sup> records a case in which menstruation ceased on January 15. The foetal movements were perceived by the mother about the beginning or middle of May, she was of enormous bulk, and expected to be confined about the 15th to the 21st October, but delivery was delayed until the 7th of December, 325 days from the cessation of the menses. The child, a male, was of much more than the usual size and weight. This was her fourth child, her first child she earned 300 days, and her second and third about 285 days. Armstrong<sup>5</sup> mentions the case of a woman who went 303 days in her second pregnancy, and 319 in her fourth. Murray<sup>6</sup> gives the following details of a woman he attended—She ceased to menstruate on February 12th, 1888, on November 28th she thought she was in labour, but the symptoms passed off. She was delivered on January 12th, 1889, of a still born child, weighing 7½ lbs and measuring 19½ inches in length, the interval between the cessation of menstruation and delivery being 330 days.

The theory that has been advanced—that protracted gestation is generally associated with unusually long inter menstrual periods—is not supported by Purkhauer's case above recorded, as the woman when not pregnant menstruated regularly every 28 days.

It will be observed that the above cases illustrate the subject of protracted gestation in a progressively increasing ratio, until the formidable total of 330 days is reached, this is not final, so far as recorded cases go, but so far as the credulity of the reader goes it probably is. In many cases of apparently protracted gestation it is more than likely that the menses were accidentally suppressed, and that impregnation subsequently occurred.

(c) **Abnormally shortened gestation** comprises the consideration of the stage of development of the fœtus at birth, and its **viability** or capacity to live after birth. There is more scope for medical evidence in shortened than in protracted gestation, in the latter the child cannot be more than fully developed, although in some cases it has surpassed the usual size, but when full term is not reached, indications of the fact are more or less obvious. In forming an opinion, too much emphasis must not be laid on the size and weight of an infant, they constitute two valuable signs of development, especially the length, which has a more fixed relation to age than the weight, but they must not be allowed to exclude other signs, nor can any reliable comparison be made between the healthy stoutness of the mother, or the converse, and the size and weight of

<sup>1</sup> *Amer Journ of Obstetrics*, 1889

<sup>2</sup> *Friedreich's Blatter f ger Med.*, 1890

<sup>3</sup> *Obstetrical Transactions*, 1885

<sup>4</sup> *Medical Times and Gazette*, 1877

<sup>5</sup> *The Lancet*, 1890

<sup>6</sup> *Brit Med Journ*, 1889

the child, as some little women habitually produce large children, and women of large stature and imposing appearance not unfrequently give birth to ill-nourished infants at full term. As in all other questions of differentiation, extremes are easy of recognition, a child at term has characteristics that clearly distinguish it from a six or seven months fœtus, on the other hand, there is little difference between an eight and a nine months fœtus, for fully a fortnight before term the child has acquired all the distinguishing characteristics of maturity. The most reliable signs of fœtal development are only available in the dead body, they are the various points of ossification, and the superficial configuration of the brain. The external indications of immaturity, in addition to size and weight, are the dark red hue and wrinkled condition of the skin, and the presence on it of lanugo, the size of the auricle, the imperfect formation of the finger-nails, the disproportionate size of the head, the absence of the testicles from the scrotum (in the male), together with general feebleness, indicated by deficiency of vigorous movements and clamorous cries with inability to suck the proffered nipple. The body-heat of immature children is subnormal, and requires conserving by means of thick layers of cotton-wool and other surroundings which lessen the loss from the surface.

**Viability.** An infant may have arrived at a sufficiently advanced period of development as to be born alive, but not to be viable— that is, not to be endowed with the capacity of continuing to live. As a general statement it may be accepted that 180 days represents the lowest limit at which an infant is viable, but it is by no means to be inferred that all infants at this period of intra-uterine life are viable. on the contrary, prolonged survival is the exception and not the rule, consequently good evidence is required to establish the occurrence of viability at this age.

Bonnar<sup>1</sup> tabulated a number of cases of premature birth tracing the after life of the infants of 22 which were born alive at 180 days of intra uterine life, only 4 lived one year and upwards, 6 more of the same group lived twenty four hours and upwards, but none of the rest longer than four months. Bailly<sup>2</sup> records the case of a child born at six months and twenty days. It was very feeble, and required much care to sustain the body heat, as an illustration of its small size it is stated that the father's finger ring could be passed over the child's foot nearly as far as the knee, thirteen days after birth the child weighed 1,250 grammes (about 2½ lbs), twelve months after birth, when the case was reported, it was living and thriving. In a well authenticated case reported by Outrepout<sup>3</sup> the child was at about 27 weeks of intra uterine life when born. It measured 13½ inches, and weighed 1½ lbs, the skin was wrinkled and covered with down, the nails were like folds of skin, and the pupillary membrane was entire, the limbs were small and were maintained in the fœtal attitude, the child was living at the age of 11 years, and had the appearance of a boy about 8 years old.

Instances are recorded in which infants born before 180 days have been reared, such cases are very exceptional and provoke scepticism as to correctness of data. Bonnar's list contains the case of one child born at 150 days which lived to the age of 19 years. A case is recorded by Moore<sup>4</sup> of a child born at the end of the fifth month, which measured 9 inches in length, and weighed 1½ lbs, at the age of 15 months the child was healthy and weighed 19 lbs. Illustrative of the risk encountered in estimating the stage of development from the size of the child is a case recorded by Hubbard<sup>5</sup>. The period of utero gestation is stated to have been about the seventh month. The infant was 10 inches long, the circumference of the head round the ears was 8 inches, round the thighs 2½ inches, the weight was 1 lb 2 ozs, the child was well formed, the finger-nails being perfect, it lived eight hours. Reference to the Table on page 26 shows that the finger nails are not perfectly formed until utero gestation has advanced eight months, but the length and weight of this child would indicate that it did not exceed six months. Barker<sup>6</sup> relates the case of a child

<sup>1</sup> *Edinburgh Med Journ*, 1865

<sup>2</sup> *Arch de Tocologie*, 1879

<sup>3</sup> *Henke's Zetschr*, 1823.

<sup>4</sup> *Philadelphia Med and Surg Reporter*, 1880

<sup>5</sup> *New York Med Journ*, 1890

<sup>6</sup> *Med Times*, 1850

born 158 days after intercourse. It measured 11 inches in length and weighed 1 lb., the nails were scarcely visible, the eyelids were closed until the second day after birth, the skin was wrinkled, three and a half years after it was thriving and healthy, but only weighed 29½ lbs. A number of similar cases have been recorded, but none of very recent date. Cullingworth<sup>1</sup> collected several, amongst which is included an instance of early viability (seventh month) that he met with in his own hospital practice.

Children born at the intra-uterine age of five months have lived for a few hours, but not much longer, it is needless to cite such cases, as it is acknowledged that a fœtus may display signs of life when born at that period of development. Without impugning the veracity of those who have reported such cases, it is nevertheless true that in no instance has absolutely convincing evidence of viability in a five months fœtus yet been recorded.

In France and Italy a child born within 180 days after marriage can be repudiated by the husband if no intercourse has taken place between him and his wife before marriage. In Germany a husband can repudiate the parentage of a child given birth to by his wife when he can prove non-intercourse from the three hundredth to the one hundred and eightieth day before the birth of the child. In Scotland a child born six months after the marriage of the mother is considered legitimate. In England and America no limit is fixed, as with protracted gestation, each case is determined on its own merits.

### SUPERFŒTATION.

The possibility of superfœtation is a disputed point amongst obstetricians, some allow that exceptionally it may occur, others regard it as being physiologically impossible. All allow the possibility of **superfecundation**—that is, the separate impregnation of two ova discharged during the same period of ovulation, the difficulty arises with respect to the possibility of an ovum derived from a subsequent ovulation being impregnated some months after the occurrence of gestation resulting from fecundation of an ovum discharged during a previous ovulation. It is now admitted on all sides that the condition of the impregnated uterus until the middle of the third month does not interpose any insurmountable obstacle to re-impregnation. The union of the decidua reflexa with the vera is not then complete, and therefore no absolute barrier exists between the ovum and the spermatozoa, the plug of viscid mucus in the cervical canal also offers no insurmountable obstacle. Those who deny the possibility of superfœtation base their opposition on the non-occurrence of ovulation during gestation, if ovulation never takes place from the commencement of gestation until after delivery, it is clear that superfœtation is impossible. The rule is that ovulation is in abeyance during pregnancy, are there any grounds for assuming the occurrence of exceptions to the rule? Galabin<sup>2</sup> instances extra-uterine pregnancy as affording evidence that ovulation may exceptionally take place during pregnancy, a five months fœtus has been found in the abdomen and one of three months in the uterus, the intra-uterine fœtus would be better situated for obtaining nourishment, and therefore its inferior development cannot be attributed to failure in this respect. The exceptional occurrence of menstruation during the early months of pregnancy does not afford more than an inferential support to the hypothesis of ovulation occurring at the same time, since menstruation and ovulation are not interdependent.

Those who deny the possibility of superfœtation explain the cases in which its occurrence has been assumed as being instances of twin pregnancy in which, as is common, one fœtus develops more than the other.

<sup>1</sup> *Obstetrical Journ. of Great Britain and Ireland*, 1878.

<sup>2</sup> *Manual of Midwifery*, 1886.

Spiegelberg,<sup>1</sup> in applying this fact, regards superfœtation as a physiologically untenable and exploded hypothesis, and considers those cases in which a fully developed child has been born, followed by another, also fully developed (within the limits attributed to superfœtation), as instances of twin fœtuses, one, much more developed than the other, being born first, and the one left behind making up for lost time and eventually appearing in as fully developed condition as the first. Cases of supposed superfœtation in which the births are more widely separated have been accounted for by the presence of a double uterus, with or without separate vagina. If only the body of the uterus is duplex, the abnormality is more likely to escape detection than when a double vagina is present. Ross<sup>2</sup> relates a case illustrative of this abnormality. On the 6th of July a woman was delivered of two fœtuses at about six months of utero gestation, on the 31st of October she was delivered of a child at full term, subsequent examination showed the existence of a double uterus. The explanation of the interval between the births of the two immature fœtuses and the child at term, is that one cavity of the uterus was prematurely delivered of twins, and that the other retained its contents up to the full period of utero gestation. In this case there was no septum in the vagina, and unless a careful examination had been made, and the cause of the abnormal course of pregnancy thus explained, another case of superfœtation would have been placed on record. A. W. Meyer,<sup>3</sup> after a review of the literature on the subject, concludes that superfœtation remains at the most only a possibility, the alleged cases being really instances of twin pregnancy with unequal development of the fœtuses.

The occurrence of ordinary twin gestation with a prolonged interval between the births is well illustrated by a case recorded by Pincott.<sup>4</sup> A woman gave birth to a weakly female infant on the 18th of October, which died the same day, after delivery the placenta came away as in ordinary confinements. On the 19th of the following November—35 days after—she gave birth to a second and finely developed female child at full term, foetal movements were felt before the membranes ruptured, but the child was still born. Postlethwaite<sup>5</sup> describes the case of a multipara who gave birth to a child weighing 6 lbs. on the 24th of February. Another child could be felt in the uterus. The mother had severe after-pains at half hourly intervals for ten hours, and after that at longer intervals for twenty four hours. She was allowed to get up at the end of a fortnight, and performed her household duties until the 4th of April, when she was delivered of a female child weighing 6½ lbs. Both children did well. Besides the long interval, 39 days, between the two births, the fact that the strong and frequent labour pains after the birth of the first child failed to dislodge the second is of interest.

Of the cases adduced in favour of superfœtation, those in which an interval of not more than from two to three months apparently elapsed between the respective conceptions, may be explained on the supposition of ordinary twin impregnation with retention of the second fœtus after expulsion of the first, even a longer interval is not impossible if the development of the fœtus which remains behind has been greatly retarded by its more favourably placed companion. If, as supposed by some, maturity of the fœtus is the initiative cause of natural labour, it is easy to see how a backward fœtus, rendered so by no inherent defect, may remain in the uterus for a considerable time beyond the limit of the normal period of gestation. The cases which, from a still longer interval between the respective conception, are regarded as proving the occurrence of superfœtation, prove too much. Towards the end of the third month the union of the decidua reflexa and vera introduces an actual barrier to reimpregnation, and a further difficulty is interposed by the then or subsequent position of the ovaries and oviducts in relation to the uterus. These obstacles, in addition to the almost invariable cessation of ovulation during pregnancy, render cases of alleged superfœtation in which an interval of five and a half months has occurred between the births of two viable children extremely difficult of acceptance. As with other negative propositions, the occurrence of superfœtation cannot well be disproved, but it may be regarded as being very close to the vanishing point of probability.

**Death of the Fœtus in Utero.**—It occasionally happens that the fœtus dies *in utero* and is retained for a more or less prolonged period, if its surroundings remain intact the access of air and of micro organisms is prevented, and either maceration or mummification of the body without putrefaction takes place. If the development of the fœtus has not advanced beyond the first month the recently formed tissues are disintegrated, so that when expelled no trace of structure is discernible. At a later period of utero-gestation there may be simply maceration of the skin, which is raised in blisters, or is entirely peeled off, the fœtus being otherwise fairly well preserved, in other cases, the fœtus undergoes mummification, and presents a flattened, dried up appearance. Schellenberg<sup>6</sup> describes a case of this kind.

<sup>1</sup> *Lehrbuch der Geburtshilfe*, 1880.

<sup>2</sup> *The Lancet*, 1871.

<sup>3</sup> *Journ Am Med Assoc*, lxxii, 1919.

<sup>4</sup> *Brit Med Journ*, 1886.

<sup>5</sup> *Ibid*, 1913.

<sup>6</sup> *Arch j Gynakol*, 1877.



A woman ceased menstruating on the 20th of June, 1876, she felt the child at the beginning of November, on the 12th of May, 1877, she was delivered of a six months fœtus, which was pressed flat and in a mummified condition, it had remained in the uterus five months after death Garrigues<sup>1</sup> describes a case occurring in a woman 37 years of age Nine months after the last menstruation a fœtus and placenta in a perfectly fresh condition were expelled, the fœtus was at about the end of the fourth month of development, the epiderm had come off, and the soft parts were atrophied In a case recorded by Hagmann,<sup>2</sup> a fœtus 16 inches long was removed by Cesarean section 441 days after the cessation of the last menstruation

The medico-legal bearing of twin deliveries, separated by an interval of two or more months, and of retention of a dead fœtus, is of theoretical rather than practical value The cases in which these conditions might come in question are when a woman gives birth to an apparently fully developed child—born alive, but dying soon after birth—within seven or eight months after death or absence of her husband, and two or three months after she is delivered of a second infant, also of full development, which lives, the heir-at-law might dispute the legitimacy of the second infant Again, if a woman whose husband had been absent for eight or nine months, gave birth to a four months fœtus eight months after her husband left her, her chastity might be called in question

The earliest period at which a woman who has been delivered of a child can again be impregnated is usually understood to be about a month after delivery, Bonnar states that it may take place as early as the fourteenth day

**Paternity and Affiliation** are determined by circumstantial evidence, and by resemblance to the father in appearance, voice, manner, gait, and other characteristics Medical evidence is directed to the minute examination of deformities, birth-marks, or other bodily peculiarities In the newly-born the questions discussed in the section on legitimacy might have to be considered

When a child is alleged to be **supposititious**, it may become the duty of a medical man to examine the child with regard to the length of time it has survived birth, and the pretended mother for signs of recent delivery If the fraud has been so successfully accomplished the suspicion is not aroused for some time after the feigned delivery (the woman having at some previous time given birth to a child at term), detection may not be easy In recent cases the condition of mother and child will probably be found not to agree, it is difficult for a woman to obtain a new-born infant at the right moment, and consequently indications of its having survived birth longer than the alleged date of confinement would reveal the fraud If the woman is examined within three or four days of the pretended parturition, the usual signs of recent delivery ought to be present When a medical man summoned to attend a woman in labour arrives after the child is born, it is his duty to assure himself that the woman has been just delivered, and that the child has been recently born, the placenta also should be examined All this is perfectly natural from the purely obstetrical standpoint and cannot be reasonably objected to Any attempts to hinder such investigations constitute grounds for suspicion, and are to be met by firm, though gentle persistence This applies to both rich and poor, with the former the legitimate succession to an estate may be imperilled, with the latter the extortion of money from an alleged seducer may be the motive In the event of absolute refusal to permit the desired examination to be made the contingencies that may arise should be explained, further objection would constitute reasonable ground for doubting the *bona-fides* of the transaction

<sup>1</sup> *American Journal of Obstetrics*, 1884

<sup>2</sup> *Monatsschr. f. Geb. u. Gyn.*, 1903

**SURVIVORSHIP.**

When two or more individuals, concerned either actually or presumptively in the succession of property, lose their lives by the same accident it may be of great importance to others concerned in the succession to determine respectively the precedence of death. In default of eye-witnesses the determination is, to a great extent, founded on conjecture, for this reason the later decisions of the courts of appeal are in favour of each case being determined by the balance of such facts as can be substantiated, rather than by inferences based on general grounds such as differences in age or sex. In the absence of conclusive evidence, two persons who perish under like conditions, and from the same cause, are assumed to have died at the same time so far as succession to property is concerned.

A number of rules have been formulated as to probabilities of the survival of one person longer than another when menaced with a common death. Age, sex, physical strength, special resources (as the capacity to swim when the cause of death is drowning) have been regarded as affording presumptive evidence sufficiently strong to justify the determination of survivorship. The possible effects of these various conditions are described when dealing with the respective modes of death.

**CHAPTER XVII****LIFE ASSURANCE.**

INSURANCE companies avail themselves of the services of medical men in two distinct capacities—as **chief medical adviser**, or as **medical examiner**. Each company has one chief adviser, and an unlimited number of medical examiners. The **chief adviser**, in addition to acting as examiner with regard to proposers who present themselves at the head office, reads and criticises all reports from the medical examiners in various parts of the country, he acts also as medical assessor or referee to the Board of Directors, and advises them as to the importance of any indications of disease, or as to the significance of details of personal or of family history. When a life cannot be regarded as first class on account of the presence of some disease, or tendency thereto, it is customary to add a certain number of years to the actual age of the applicant and to demand a correspondingly increased annual payment, under such circumstances the chief medical adviser has to estimate the probable duration of life, and to fix the age at which the proposal shall be accepted. He has to satisfy himself that there are no omissions nor obvious errors in the examiners' reports, and if he detects any discrepancy his duty is either directly or through the office to communicate with the examiner, and to request an explanation.

**Medical Examiners** are selected by the actuary or other officer of the company in those towns in which an agency is established, in large towns all the proposers are usually sent to one medical man, who in this way becomes an adviser of the company, in country places any medical man may be selected whose residence happens to be conveniently near to that of the proposer. When the policy is for a very large sum, the office may require the proposer to be examined by two medical men. The object evidently is to secure as far as

possible, a thorough investigation of the physical condition and functional activity of the various organs of the body, to accomplish this the two medical men should make the examinations separately, an arrangement, however, which is usually not specified, and as competition between rival offices is very keen, the agents favour a conjoint examination by the medical men in order to avoid deterring the applicant by the prospect of having to undergo two distinct examinations.

Insurance offices vary in their requirements as to reports—some provide elaborately arranged printed forms which suggest all the necessary investigations, and contain a long string of questions relating to personal and family history to be answered by the applicant, which are severally asked, and the answers recorded by the medical man at the time the examination takes place. Printed forms of exhaustive questions have one distinct advantage, their routine character enables questions of a more or less delicate nature to be put without causing offence, this applies especially to syphilis, the possibility of the presence of which in a latent form is not to be lost sight of. Other offices, in the letter to the medical man requesting him to examine an applicant, direct attention to the chief subjects on which information is required and ask him to furnish a report in the form of a letter to the actuary—when this is the case the applicant himself fills in his statement of family and personal history and sends it direct to the office.

All such statements must be absolutely correct, an inaccuracy, even though not intentional, may vitiate the contract. Amongst other questions the applicant is asked whether he has been refused by other offices. Any attempt to evade giving a direct answer to this question is fatal. In one instance an applicant stated that he was still corresponding with other offices, but omitted to mention that he had been declined by offices previously applied to, another merely stated that he was insured in other offices, but did not mention that he had been declined by several, in both these cases the courts cancelled the contract (*Lond Ass Co v Mansel*, 11 Ch D 370, *Gen Pror Ass Co ex parte Duintree*, W R 1870). The question, "Who is your ordinary medical attendant?" includes the person who may be actually in attendance at the time, although he may not be the ordinary attendant; on the other hand, it is not sufficient to give the name of a medical man who was last casually consulted, and to omit that of the usual medical attendant, the object of the question being to ascertain the name of the medical attendant that is best able to give an opinion as to the applicant's state of health (*Everitt v Desborough*, 5 Bng 503). If at the time of making the statement the applicant truthfully says that he has never had a serious illness, and subsequently, but before the policy is effected, is taken seriously ill, and does not inform the office, the contract is void (*Canning v Farquhar*, L R, Q B D 1886). Some allowance, however, is made for excusable want of knowledge in interpreting symptoms indicative of disease, for instance, an applicant stated, in answer to inquiry, that he had never had gout, although really he had suffered from certain symptoms of that disease without knowing it, the court ruled that as only a medical man could have detected the nature of the symptoms, the applicant could not reasonably be expected to have done so (*Fowkes v Manch and Lond Ass Co* 3 F & F 440). The exact meaning and interpretation of other questions put to the applicant has been the subject of judicial ruling. Spitting of blood has been held to mean such as is due to a disease which tends to shorten life, and not such as may arise from accidental cause, or from temporary ulceration of the mouth, in the same way a cough means one of constitutional origin, and not a temporary ailment. Again, a declaration by an applicant that he had not been afflicted with nor was subject to fits, was held to mean not that he had never had a fit, but that he was not habitually or constitutionally afflicted with fits. In this case the applicant, in consequence of a fall, had had two epileptic fits, within a short interval, several years before the date of the policy, but which the jury were satisfied had never recurred (*Chattock v Shaw*, 1 M & R 498, *Shilling v The Acc Death Ass Co*, 1 F & F 116).

No one should be present at the examination except the applicant and the medical examiner. The first thing is to observe the general appearance of the applicant—the complexion, the healthy or cachectic colour of the skin, the

presence of enlarged veins on the nose or cheeks, puffiness or redness about the eyes, together with the general expression, whether it affords indications of latent physical disease, or of mental anxiety. The gait of the applicant as regards spasticity, ataxia, or other deviation from the normal, and the state of the tongue, gums, teeth and throat are to be noted. Inquiries are to be made as to the presence of hernia and other pathological conditions. The height, the weight, and the measurements of the chest at the level of the fourth rib, during full inspiration and deep expiration, should be ascertained. The expansion of the chest should also be inspected and the respiration-rate taken, this, and the rate of the pulse, which should also be counted, is frequently influenced by nervousness of the applicant. The lungs and the heart are to be examined in the usual way (the skin being bared) and the result recorded. Special attention should be devoted to the apices of the lungs, back and front, especially if the family history is indicative of phthisis. The character of the breath-sounds as regards harshness, vesicular breathing, prolongation of the expiratory sound, and lessened audibility of the inspiratory sound, together with increased vocal resonance at the apices, are all of significance. If a heart murmur is present, the valves implicated and the character of the murmur are to be recorded, if the murmur is mitral-regurgitant it is well to ascertain how far posteriorly it can be heard, as affording some clue to the amount of regurgitation and consequently of the importance of the lesion. In some nervous people a sound resembling that of a murmur or organic origin may be heard on applying the stethoscope, such a sound disappears after the first impression produced on the nervous system by the momentous nature of the examination has passed off. It is to be borne in mind that even strong healthy men are often influenced to an extraordinary degree in this way, the very fact of not having undergone medical examination before, with the possibility presented to their minds that some deadly latent disease may be discovered, produces no little mental perturbation. If there is a gouty family or personal history the tension of the pulse and the daily amount and the colour of the urine should be ascertained. In all cases a specimen of urine voided in the presence of the examiner, or with such precautions as preclude the possibility of deception, should be examined as to naked-eye appearance, reaction, specific gravity, and the presence or absence of albumen, if albumen is present a microscopical examination should be made for casts. In the case of applicants of or above middle age, especially if the urine is pale and of low specific gravity, more than a cursory examination for albumen should be made. As is well known, the urine from cases of granular kidney is light coloured, of low specific gravity, and often contains a mere trace of albumen, and at times none at all, such urine examined off-hand, with nitric acid, may appear free although a little may be present, after adding the acid, the specimen should be allowed to stand for five or ten minutes, and then examined. It is preferable to nearly fill a test-tube with the urine and to boil the upper stratum, a slight haziness, where the heat has been applied, which persists after the addition of a few drops of acetic acid, reveals the presence of a very small amount of albumen. Urine that has an alkaline reaction should be rendered *slightly* acid by the addition of a drop or two of acetic acid *before* boiling. If the specific gravity is above 1,020 the urine should be examined for sugar, and in any case if there are reasons for suspecting diabetes. Discussion as to the probable duration of life when albumen is present in the urine, or when other abnormal conditions exist, is beyond the scope of this book.

Insurance companies not unfrequently send a separate form to the usual

medical attendant of the applicant requesting answers to the questions thereon printed. It is optional on the part of the medical attendant whether he answers the questions or not, but if he does so and accepts the fee, he is bound to answer fully and without reserve. Some offices ask the applicant's permission to interrogate his medical attendant, if given it is supposed in this particular instance to relieve the attendant from any obligation to professional secrecy. Although the office promises to treat the answers of a medical referee as confidential, the document is not a privileged communication, it must be produced, if required, should subsequent proceedings be taken, the same applies to the answers of a lay referee (*Mahoney v Nat Widows' Fund*, 19 W K 722). The medical referee cannot recover any fee for information given unless there is an agreement to that effect between him and the office.

Formerly it was customary for insurance offices to bar death from suicide, a restriction which gave rise to medico-legal discussions in relation to insanity as a cause of the suicidal act. It has been decided that the word "suicide" means *homicidium sui ipsius*—death by his own hands—and not merely felonious suicide. "It includes every case of self-destruction by a man knowing the probable consequences of his act, and doing the act voluntarily and intending the consequences to follow" (*Clift v Schwabe*, 3 C B 437). At the present time it is the custom to take the risk of suicide along with other risks, provided that death from this cause does not occur within a certain period, which has been fixed by most offices at one year after the payment of the first premium—that is to say, after the first renewal of the policy, by the payment of the premium due at the expiration of one year from the date on which the policy was granted, the risk of suicide is accepted by the office, and in the event of its taking place, the sum insured for is paid. Some offices retain older customs, and simply return to the legal representatives of the holder of a policy who has committed suicide the money which has been paid in premiums.

Information as to personal and family history to be furnished by the applicant —

Name, age, occupation, civil state. Have you ever had any of the following diseases? Dizziness, unconsciousness, epilepsy or convulsions of any sort, apoplexy, paralysis, or any disease of the nervous system, severe or persistent headache, discharge from the ears, chronic cough or hoarseness, asthma or other chest diseases, spitting of blood, disease or functional disturbance of the heart, dyspepsia, chronic diarrhoea, colic—renal, hepatic, or intestinal, piles, fistula or other diseases of the rectum, bladder or kidney disease, stricture, syphilis or venereal diseases, malarial or other fever, rheumatism or gout, cancer, tumour or ulcer, or chronic skin disease. State the full particulars of any other illness, disease, or injury you have had. Has your weight recently increased or diminished? When did you last consult a medical man? State your habits, past and present, as to the use of alcohol, tobacco, and narcotics. Have you ever been under treatment in an asylum or refuge? Are you now in good health so far as you know and believe?

The family history includes the ages of the applicant's father and mother and of their parents, and of the applicant's brothers and sisters, with information as to their respective states of health. If dead, the age at the time of death and the cause of death, with the duration of fatal illness, are demanded.

#### FORM OF MEDICAL REPORT IN LIFE ASSURANCE

Figure and general appearance ?	During deep expiration ?
Apparent age ?	Number of respirations per minute ?
Weight ?                      Height ?	Is there anything abnormal in the character
Measurement of abdomen at the level of umbilicus ?	of the respirations ?
Of the chest, at the level of fourth rib, during full inspiration ?	Is there any indication of disease, either acute, or chronic, of the respiratory organs ?

State the rate and tension of the pulse ?

Is it intermittent or regular ?

Is there any indication of disease of the heart or blood vessels ?

Test the urine, voided at the time of examination, and state

Clear or turbid ?      Colour ?

Specific gravity ?      Reaction ?

Presence of albumen ?      Of sugar ?

Result of examination by the microscope ?

Is the applicant personally known to you, and if so, how long ?

Is there any bodily malformation, impairment of sight or hearing, or loss of any part of any member ?

Has the applicant been successfully vaccinated ?

Has he a hernia ?

Is the risk affected by anything in his residence or occupation ?

Are you satisfied that there is nothing in his physical condition, habits, personal or family history not distinctly set forth tending to shorten his life ?

Do you unqualifiedly recommend the applicant for insurance ?

The burden of the proof of death rests with the representatives of the insured, who must tender satisfactory evidence to the insurers (*Strong v Harvey*, 3 Bing 304, *Braunstein v Acc Death Ass Co*, 1 B & S, 782). In ordinary cases a certified copy of entry of burial in the general or the district registry is required, and usually an additional certificate, signed by the medical attendant of the deceased, stating the cause of death and the duration of the illness.

### ASSURANCE AGAINST ACCIDENTS.

The intention of accidental assurance companies is to insure against injuries and death of accidental causation only, and therefore their policies usually contain words to the effect that the assurance does not extend to injury arising from natural disease, weakness, or exhaustion consequent on disease, disease caused by injury, and injury caused by disease, are usually debarred. The interpretation of the bearing of these limitations, as regards contentious cases, is a fruitful source of litigation.

For example, a man assured against accident had an epileptic attack while fording a stream and was drowned. The office declined to pay the insurance on the ground that the fit—i.e., disease—was the actual cause of death, the Court decided that as death was due to drowning, it was of the nature of an accident though caused by the fit. In another similar case, a man, while standing on the platform of a railway station, had a fit, he fell across the line and was killed by a train, in this case also the office was held to be liable (*Lawrence v Acc Assur Co*, L R, Q B D 1881). In a case in which the exceptions included "death from hernia or any other disease, whether causing death directly, or conjointly with accidental injury," the insured accidentally fell on the floor and ruptured himself, he died shortly after from the effects of the rupture and a surgical operation necessary to relieve it, it was decided that death was due to accident and not to disease. (*Filton v Acc Death Assur Co*, 17 C B, NS, 122). In another case, however, in which after "or any other disease," the words "or secondary cause or causes," were introduced in the policy, the insured having died from erysipelas which supervened after he had accidentally cut his foot with a piece of pot, the court held the office protected. The Chief Baron, in dissenting, held that not only the actual injury but any disease, as lockjaw, hemorrhage, or erysipelas caused by the injury, is covered by the policy (*Smith v Acc D Assur Co*, L R, 5 Ex 302). In the following instance the limiting clauses were much more liberally interpreted. The insured fell and dislocated his shoulder, he was put to bed, but, from exposure of the body due to restlessness, he developed pneumonia and died in a month after the accident, having been confined to bed all the time, the Court decided that death was caused by the accident within the meaning of the policy (L R Q B D 1889). A common clause in the policy exempts the office from liability for injury happening by exposure of insured to obvious risk of injury, as when the insured crossed a railway in daylight, at a place not intended as a crossing, and was killed by a train. The Court decided that the office was not liable (*Cornish v Acc Assur Co*, 23 Q B D 453).

**The Workmen's Compensation Act, 1906.**—Under this Act employers are liable to pay compensation to workmen who sustain "personal injury by accident arising out of and in the course of the employment" If the workman is killed, compensation is payable to the dependants.

The term "Workman" as defined by the Act "does not include any person employed otherwise than by way of manual labour whose remuneration exceeds £250 a year, or a person whose employment is of a casual nature and who is employed otherwise than for the employer's trade or business, or a member of a police force, or an outworker, or a member of the employer's family dwelling in his house, but save as aforesaid means any person who has entered into or works under a contract of service or apprenticeship with an employer whether by way of manual labour, clerical work or otherwise, and whether the contract is expressed or implied is oral or in writing."

The Act also provides that persons suffering from certain specified diseases due to the nature of their employment shall be entitled to compensation just as if they had incurred personal injury by accident. The more important of these conditions are anthrax, poisoning by lead, mercury, phosphorus, or arsenic, glanders, ankylostomiasis, compressed air illness, miners' "bent hand" and "bent knee," miners' nystagmus, writer's cramp, telegraphist's cramp, and dermatitis, or ulceration of the skin, or ulceration of the mucous membrane of the mouth or nose, produced by dust or liquids. A certificate that a person is suffering from a disease to which the Act applies is given by a certifying surgeon appointed under the Factory and Workshop Act, 1901, in accordance with rules made under the Workmen's Compensation Act.

Medical men may also be appointed Medical Referees under the Act, their duties being to determine or report upon various medical questions. Thus if an employer or workman is aggrieved by the refusal of a certifying surgeon to give a certificate of disablement or to suspend a workman from his employment on the ground that he is suffering from a scheduled disease, the matter is referred to a medical referee, and his decision is final. In a case where after a medical examination an employer and a workman are unable to agree as to the workman's condition or fitness for employment, if both parties apply to the Court the matter may be referred to a medical referee. The latter examines the workman and gives a certificate as to his condition and fitness for employment. This certificate is conclusive evidence as to the matters so certified. Other duties are to report upon medical questions arising in the course of an arbitration, and to examine and certify as to the condition of persons in receipt of compensation who propose to live outside the United Kingdom, such persons being only entitled to compensation if the incapacity is certified to be permanent. Finally in cases presenting difficulties of a medical character, a county court judge may summon a medical referee to sit with him as assessor.

The Workmen's Compensation Act has given rise to an immense amount of litigation, particularly in reference to the definition of "accident", the interpretation of the words "arising out of and in the course of the employment", and the complication introduced by pre-existing disease, or disease following and attributed to an accident. The decisions have frequently turned upon points of considerable subtlety, and to cite cases without quoting the arguments used at some length might be misleading. For recorded cases and decisions some of the numerous special books on the subject should be consulted.

### MALINGERING.

**Malingering** in the strict sense is the fraudulent imitation or exaggeration of injury or disease for the purpose of obtaining some benefit, or of avoiding work. But the term is also somewhat loosely applied to a class of persons who, while suffering from some injury or affection not of a serious character, have unconsciously exaggerated their symptoms until they are convinced that they have a just claim for compensation or sickness benefit. The question became of importance with the passing of the Workmen's Compensation Act in 1906, and has assumed greater prominence since the National Insurance Act came into force in 1912.

The **real fraudulent malingerer** is relatively rare in comparison with the great number of claims which are preferred under the two Acts mentioned above. But in the extensive literature devoted to this subject which has now grown up, a large number of fraudulent claims, many of them of a highly ingenious character, are described. Four types of cases may be recognised - (1) Persons who have no affection or injury of any kind, (2) persons who are suffering from lesions deliberately produced, (3) persons who represent old-standing disease or injury to be the result of a recent accident, (4) persons who are suffering from the effects of an injury or disease not of a serious character, and who consciously and fraudulently exaggerate their symptoms.

As instances in the first group may be cited persons who allege that they are suffering from stiff or deformed joints resulting from a fall or other accident. The knee is frequently chosen. A person who is practising this fraud will limp painfully into the consulting-room, cry out when the joint is examined, and forcibly resist attempts to flex it beyond a certain angle. An X-ray examination will fail to reveal any pathological condition. Sometimes the fraud may be detected by carefully watching the person when he thinks he is unobserved. It may be noticed after the examination is over, and while the doctor is apparently busy writing his notes, that the individual in order to put on his boots now readily bends a joint which has just resisted all attempts to flex it. Pain and stiffness in the back is another condition frequently simulated. In these cases the patient should be asked to indicate exactly the spots at which the pain is felt. On going over the areas again at the end of the examination, or at subsequent examinations, a different set of areas may now be described as painful. Sometimes the electric battery may be usefully employed. Collie describes the case of a man who complained loudly of pain when touched in the dorsal region of the back. He had been told that it was characteristic of the particular condition from which he suffered, that no pain was caused by an electric battery. When the current was applied he stated that he felt nothing, and when the strength was increased he bore it manfully for a time, but at length collapsed with a howl and admitted that he had been practising an imposture. As with joints of the limbs, fraudulent assumption of stiffness in the back may be detected by watching the man while off his guard, a rigid back often becoming suddenly supple as he stoops to draw up his trousers.

Patmore relates a case in which elaborate precautions were adopted to prevent examination. A medical student encased his leg in plaster of Paris and then made claims on several insurance companies. He was visited by various medical men, but they did not disturb the casing, and it was not until one, whose suspicions were aroused, insisted upon removing the plaster, that the fraud was discovered.



Sometimes foreign substances are added to the urine. Morley<sup>1</sup> met with a case of a man who was supposed to have had an injury to the pelvis, and who produced while in hospital several specimens of dark-coloured urine, looking as though it was blood-stained. No corpuscles could be detected under the microscope, and specimens drawn off with a catheter was found to be normal. The man was kept under observation, and was discovered to have a supply of saffron threads which he had introduced into his urine.

Affections of the special senses, such as partial or complete blindness or deafness, may be feigned. A person may allege that he is unable to read test-type, but if given a glass which he believes will help him, the vision will suddenly improve although the glass be a *plain* one. In another case a man professed not to be able to see a white object placed on the dark floor, but when after an examination of his chest he missed a black safety-pin which the doctor had deliberately pushed from the table on to the floor, he instantly stooped down and picked it up.

Feigned deafness may be difficult to detect. If limited to one ear, by alternately plugging the ears, testing the hearing, and removing the plugs, the man may become confused, and may be found to be hearing quite well with the sound ear plugged and the affected ear free.

As instances of real injuries deliberately inflicted in order to support claims may be mentioned skin lesions produced by the application of strong acids and other irritants. Such lesions are usually found on the front of the body, and if the person be kept under observation will show anomalies and variations in healing which will enable them to be differentiated from known skin diseases. Norman Walker<sup>2</sup> describes the case of a girl upon whose arms and hands eruptions continued to appear for months. They healed up under treatment, only to break out elsewhere, until there had been a complete renewal of all the skin on the fingers. Fraud being suspected, the girl was removed to a hospital and watched, but although her bed and clothes were frequently searched, nothing was found, and the eruptions continued to appear. At length, a small bottle containing carbolic acid was found concealed in her handkerchief and it appeared that whenever she left her bed she had managed to hide it in this way. The eruptions rapidly healed after the bottle was removed. This case was probably of an hysterical character.

Irritation of the eyes is sometimes practised until keratitis or perhaps panophthalmitis is produced.

Mutilation of fingers or toes is not infrequent. An extreme case occurred in Wales, in 1905. A man claimed damages for injury against a railway company, his story being that while travelling in a train he was thrown by a violent jerk against the door, which flew open and allowed him to fall out. He lost consciousness, but recovered as another train was passing over his legs, which in consequence had to be amputated below the knee. For the company it was shown that the plaintiff had previously purchased a number of newspapers containing insurance coupons, that there were no marks of injury attributable to the fall except an abrasion over one eye, although he weighed over 15 stone and the train was travelling at 28 miles an hour, that he had told several persons he had dreamed he would lose both legs below the knee, that when found on the line he showed no signs of recent unconsciousness, and that all the doors were found properly closed when the train arrived at the station next to the place where the accident was alleged to have occurred. A verdict was found for the company.

<sup>1</sup> Int. Congr. of Med., 1913

<sup>2</sup> Brit. Med. Journ., 1910

Diseases which are fraudulently represented as having been caused by accident are, most frequently, those affecting muscles or joints, such as lumbago, gout arthritis, rheumatism, etc. An X-ray examination will often give great assistance in coming to a conclusion. Enlarged glands or buboes in the groin are sometimes alleged to have been caused by a strain.

Injuries to the fingers or hands leading to some degree of stiffness of a joint are frequently exaggerated and alleged to have caused total incapacity for work. In many of these cases the condition could be cured or ameliorated by massage or passive movements, but the claimant refuses to undergo treatment. Another type of injury is that in which a nerve is involved in the stump of a severed finger, producing pain on pressure and preventing the handling of objects. A slight operation would set many of these cases right, but the courts will not insist upon an operation unless it can be shown that it is devoid of risk and will remove the disability.

The class of persons who present claims which are not justified, but who nevertheless are not malingerers in the true sense of the word, inasmuch as they honestly believe their claims to be genuine is much larger and gives rise to the great bulk of the cases which come before the law courts. In a typical case of this sort a person has met with an accident, not of a serious character, from the effects of which he would under ordinary circumstances rapidly recover. But in consequence of injudicious sympathy, of hearing experiences of other workmen who have received compensation, and of the prospect of so benefiting himself, he is led to magnify both the accident and the results thereof. He makes a claim and is surprised to find that others do not regard his injuries as being of so serious a character as he does himself, and thus a sense of unjust treatment is added to his mental impressions. He may be advised to persevere in his claim by persons who are not always disinterested in the matter. These cases frequently arise in connection with injuries to the joints, muscles, or back, and there is every gradation between them and the true neurasthenic or hysterical individual with functional paralysis and loss of sensation.

An important sub-group comprises persons who in consequence of an accident undergo a medical examination, and then for the first time pre-existing disease is discovered. A person gets a splinter into his eye, for example, and on examination it is found that he is suffering from senile cataract. He was previously quite unaware of the affection, and usually nothing will convince him that the condition must have been coming on for a long time and is not the result of the accident. In other cases general paralysis of the insane, tabes dorsalis, heart disease, or aneurism is discovered.

## CHAPTER XVIII

## MEDICO-LEGAL BEARINGS OF DIVORCE.

MARRIAGE is of the nature of a contract and is not legally binding if one of the parties to it is incapable of consenting to, or fulfilling, the contract at the time it was entered into. In relation to divorce there are two questions which come within the province of the medical jurist, the first is **unsoundness of mind** preventing consent this constitutes a civil disability which invalidates marriage, the second is **impotency** or **physical incapacity** for sexual intercourse this constitutes a canonical impediment to marriage.

The plea of **insanity** can only be offered in a suit for nullity of marriage when the mental disorder existed at the time the marriage took place, the fact that one of the contracting parties was insane at the time the contract was entered into renders the contract null and void, because insanity incapacitates the person subject to it from giving a rational assent to the terms of the contract. Insanity rarely develops without warning, but salient symptoms may suddenly show themselves after the mind has been some time diseased, the previous condition not being sufficiently marked as to give rise to suspicion of mental disorder, marriage, in such cases, not unfrequently kindles up the dormant state of mental disease into one of an acute character. In some instances the relatives and friends of the affected party conceal the disorder from the person with whom the marriage contract is to be made, with the forlorn hope that the new domestic relations will dissipate the existing trouble, the result almost invariably being disastrous to all concerned. When insanity is pleaded as a cause for nullity the diseased mental state is usually observed shortly after the marriage takes place. In the case of *Hunter v. Edney* (Divorce Court 1882) the wife refused to allow the marriage to be consummated; and an investigation of her mental condition showed that she was suffering from melancholia, and probably had been for some time, in granting a decree, the judge pointed out that the woman did not appear capable of understanding actions free from the influence of delusions, and was, therefore, incapable of entering into a contract like that of marriage.

The plea of insanity in relation to proceedings in the divorce court took a new departure in the case of *Hanbury v. Hanbury* (Divorce Court, 1892). The wife brought an action against the husband for dissolution of marriage because of his adultery and cruelty, the defence was that the respondent was suffering from insanity when the acts complained of were committed. The President, Sir C. Butt, in summing up said that it had been argued that insanity was an answer in a suit for dissolution of marriage, and he was far from asserting that it was not in any case, but he thought to make the plea a good one, the insanity must be lasting. It might be that the plea would be a good one if the insanity of the person were such as to necessitate his being placed in an asylum, from which there was no prospect of his discharge because there was no prospect of his recovery, or of amelioration of his malady. A decree *nisi* was granted, and from this it seems that insanity might be admitted as a plea for irresponsibility. The case in point did not decide the question,

because the respondent had given way to great excess in alcohol, a fact that weighed with the jury and confused the issue so far as precedent in relation to insanity itself is concerned

In order that incapacity to take part in the performance of sexual intercourse can be regarded by the Court as a ground for a decree of nullity, it must be proved that the defect existed at the time of marriage and that it is of a permanent nature. It is not essential that the malformation should be of such a nature as absolutely to prevent coitus, if it interposes only a partial obstacle it constitutes a ground for a suit of nullity. The incapacity must be permanent, if a cure is at all possible, though it may be extremely improbable, it is sufficient reason for refusal of a decree, if, however, an operation to effect a cure would be attended with great danger to life, the malformation is regarded as incurable. If the impediment did not exist at the time of marriage, but has occurred since, it does not constitute a ground for nullity.

In cases where there is no apparent physical defect a cohabitation of three years' duration is required to ascertain with whom the fault lies. This rule is not invariable. If the Court deems that a sufficient time has elapsed for the difficulty to be overcome were it not of a permanent nature, a decree may be granted sooner, in one case, upon an apparently conclusive medical certificate that the impotence would be permanent, nullity was pronounced after a cohabitation of three months only. Triennial cohabitation is not required when the incapacity of husband or wife is of a visible and incurable nature which can be ascertained at once.

Wilful refusal of marital intercourse is not sufficient to obtain a decree, although persistent resistance on the part of the wife has been so regarded. Repeated attacks of hysteria in the wife brought on by the husband's attempts at intercourse has, after a cohabitation of three years without consummation, been deemed ground for a decree *nisi*.

Impotence does not render a marriage void, but only voidable—that is, the injured person must take active proceedings and sue for a decree. If sufficient grounds have existed, such as an absolute physical impediment to intercourse so that the marriage has not been consummated, and no proceedings are taken during the lifetime of husband and wife, the marriage cannot be declared void after the death of one of them, this has been attempted in order to deprive the husband of benefits from a wife who died intestate, the next of kin claiming the estate.

It is incumbent on the petitioner for a decree of nullity of marriage on the ground of impotence to prove that marriage has not been consummated. The necessary evidence is obtained by medical examination of the generative organs of the respondent, which is made by two medical inspectors appointed by the Court. The husband and wife may conjointly select two medical men, or if they cannot agree each may nominate one, the petitioner then moves the Court for the appointment of the medical men chosen as inspectors, and at the same time moves for an order that the respondent submit to the inspection. The two medical inspectors after appointment have the following oath administered to them, which sufficiently explains the duties they are called upon to fulfil—

"In the High Court of Justice Probate, Divorce, and Admiralty Division (Divorce).

"A B and C D, doctors of medicine

"You are produced as inspectors in a cause depending in the Probate, Divorce, and Admiralty Division of the High Court of Justice (Divorce) entitled . . . falsely called . . . , to examine the parts and organs of generation of . . . ,

the petitioner in this cause, and also of \_\_\_\_\_, falsely called \_\_\_\_\_ the respondent in this cause

" You respectively swear that you will faithfully, and to the best of your skill, inspect the parts and organs of generation of each of them, the said \_\_\_\_\_ and \_\_\_\_\_, and make a just and true report in writing, to the Right Honourable the President of the above Division, whether the said \_\_\_\_\_ (*the petitioner*) is capable of performing the act of generation, and if incapable, whether such, his incapacity, can be cured by art or skill, and also, whether the said \_\_\_\_\_ is or is not a virgin, and whether she hath or hath not any impediment on her part to prevent the consummation of marriage, and that one of you will deliver such report under your hands and seals, closely sealed up, to one of the registrars of the above Division

<p>" Sworn at the Principal Registry of the Probate, Divorce, and Admiralty Division of the High Court of Justice, this _____ day of _____ 19 _____, Before me, _____ _____, Registrar "</p>	}	<p>(Signatures of the medical inspectors )</p>
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Medical evidence will be received upon the question of incapacity, although no application has been made for an order for personal inspection of either of the parties

## CHAPTER XIX

### MODES OF DEATH RESULTING CHIEFLY FROM ASPHYXIA.

#### HANGING.

DEATH from hanging is produced when the body is wholly or partially suspended by means of a cord or other ligature round the neck, until life is extinct. It is not necessary that the body should be in the upright posture, nor that it should cease to rest on the ground or other means of support, as will be presently shown, a very slight degree of tension on the ligature, such as that caused by partial suspension, is sufficient to produce death. Apart from judicial hanging, the post-mortem appearances are those of death from **asphyxia**, together with indications of the manner in which asphyxia was produced.

In by far the greatest number of cases of death from hanging the **ligature** surrounds the neck **above** the **thyroid cartilage**. Even if the noose is originally placed lower down, it is not in the first instance drawn tight enough to prevent the weight of the body dragging the neck as far through it as it can come. In 153 cases observed by Maschka<sup>1</sup> the ligature was above the thyroid cartilage in 149, on it in 1, and below it in 3. When the ligature is above the thyroid cartilage slight constrictive force suffices to occlude the air-passages. Langreuter<sup>2</sup> investigated on the dead body the mechanism of the closure of the air-passages by removing the skull-cap and brain in the ordinary manner, and cutting away the base of the skull so that the laryngo-pharyngeal region came into view, the subjects of the experiments had died from natural causes—they were not cases of death from strangulation. A cord, placed round the throat, between the thyroid cartilage and the hyoid bone, was carried up under

<sup>1</sup> *Vierteljahrschr f ger Med*, 1886.

<sup>2</sup> *Handbuch*, Bd 3

the angle of the lower jaw. By moderate traction in the long axis of the body, as in hanging, the epiglottis was pushed against the back of the pharynx, by stronger traction the base of the tongue followed, and the free end of the epiglottis was pressed between the base of the tongue and the posterior pharyngeal wall, only moderate force was necessary to close entirely the air-passages. If the cord was placed below the thyroid cartilage, or horizontally upon it, and traction made as before, it slipped up above the cartilage, if retained on the cartilage with the hand, the vocal cords could not be completely approximated even when great force was used in pulling the constricting medium upwards and backwards. On suspending the body, the cord at once slipped up immediately under the lower jaw, and on looking from above, the tongue was seen to be displaced, upwards and backwards, and pressed into the opening in the base

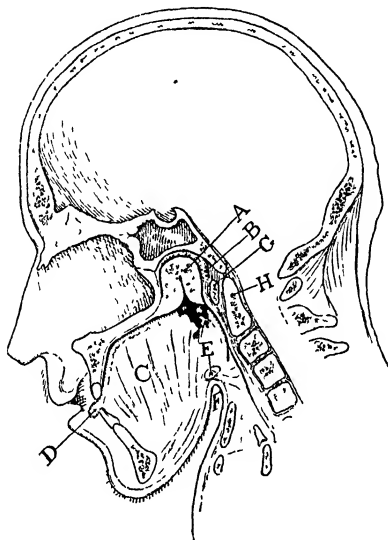


Fig 15—A, soft palate, B, wall of the pharynx, C, tongue, D, tip of the tongue pressed between the teeth, E, body of the hyoid bone, F, groove made by the rope, G, anterior portion of the atlas, H, odontoid process of the axis.

of the skull. Most of the experiments were repeated with a rolled-up handkerchief, the results were the same, but more force was required. Ecker<sup>1</sup> demonstrated like results with the body of a man which was found hanging from a tree in winter, frozen so hard as to be easily sawn into vertical sections. The soft structures at the posterior part of the floor of the mouth were doubled up into the cavity of the pharynx so as completely to fill and obliterate the naso-pharyngeal passage. Fig 15 is an exact representation of what was found.

When hanging causes complete occlusion of the air-passages, death results

<sup>1</sup> Virchow's Arch, 1870

for the most part from asphyxia. Another factor, however, has to be considered—compression of the large blood-vessels of the neck which may produce effects equal to or even exceeding in importance those due to asphyxia, in any case these vessels will undergo compression, and, if the air-way escapes complete occlusion, arrest of the intra-cranial circulation may be the principal cause of death. When death begins in the brain, as the result of compression of the large vessels of the neck, it is not so much from excess of intra-cranial blood-pressure, as from arrest of the cerebral circulation, it is very rarely that cerebral hæmorrhage takes place. Illustrative of this mode of death is a most exceptional case, recorded by Reineboth,<sup>1</sup> of a man who hanged himself about two months after tracheotomy had been performed on him for malignant disease of the throat, the canula being left in the trachea, the noose of the cord surrounded the neck above the canula and thus left the air-way to the lungs quite free. At the necropsy the arteries of the base of the brain and the substance of the pons and medulla were gorged with blood, the rest of the brain-substance and the vessels of the pia contained but little blood, no sub-pleural ecchymoses nor other indications of asphyxial death were present in the thorax. If the air-way escapes occlusion, or is only partially occluded, a much longer time elapses before death takes place than if respiration is completely arrested, this has been shown experimentally with a dog: an opening was made in the lower part of the trachea, and the animal was hung by the neck with a cord, the noose being above the opening, a much longer time was required to cause death than is the case in ordinary hanging, and the post-mortem signs are different.

Great importance was formerly attributed to the respective influence of pure asphyxia and of what was called "apoplexy" as proximate causes of death from hanging. This resulted from the inconstant post-mortem appearances presented by the brain and by the lungs: sometimes the brain, sometimes the lungs, and sometimes both are found gorged with blood. It was inferred that if death took place from asphyxia the post-mortem signs ought to be invariable, as they were found not to be invariable, hypotheses were formulated to explain the discrepancy. It was assumed that death from hanging might be brought about in two distinct ways—by pure asphyxia and by apoplexy without asphyxia, seeing, however, that in a large number of deaths from hanging both lungs and brain are hyperæmic, a mixture of the two causes of death was described to suit this post-mortem condition. In consequence of these views the post-mortem appearances respectively produced by two closely allied modes of death were frequently treated as though they resulted from separate and distinct causes, and variable external conditions were regarded as necessary factors in the production of these dissimilar results.

Experiments, however, show that the preponderance of the signs of death from asphyxia over those indicating disturbance of the intra-cranial circulation, or the converse, are not solely dependent on the varying degrees of compression produced by the ligature on the air- and blood-channels respectively. Precisely similar external conditions may cause, in one case excess of blood in the lungs but not in the brain, and in another excess of blood in the brain but not in the lungs. It is probable that the state of the lungs, as regards inflation at the moment the air-passage is occluded, has a good deal to do with the relative hyperæmia of brain and lungs, some experiments made by Patenko<sup>2</sup> tend to prove this. A number of dogs were hanged, some at the end of full inspiration, others at the end of full expiration, and the distribution of the blood in the

<sup>1</sup> *Vierteiljahrsschr. f. ger. Med.*, 1895

<sup>2</sup> *Annales d'Hygiène*, 1885

various internal organs was observed after death. In those hanged at the end of full *inspiration* the lungs contained little blood, but the sinuses of the skull, the membranes of the brain, and also the vessels of the abdominal organs, contained a great deal. In the animals hanged at the end of full *expiration* the lungs contained much blood, and the cephalic and abdominal vessels little. When the air-passages are occluded at the end of full *expiration* the blood flows from the periphery into the heart and—on account of the negative pressure developed in the thoracic cavity—from thence to the lungs, it cannot leave the lungs on account of the damming up that occurs in the dilated pulmonary vessels and from the want of intra-pulmonary pressure. On the other hand, when occlusion takes place at the end of full *inspiration*, the pressure on the lungs causes expulsion of the blood they contain and consequent excess in the cranium and in the abdominal vessels. Hofmann<sup>1</sup> considers that retraction of the abdominal walls, especially at the pit of the stomach—which occurs simultaneously with the efforts at respiration—compresses the lungs, and thus drives out the blood from them.

Strong corroborative evidence that the want of uniformity in post-mortem appearances of death from hanging is not dependent (or is so only under exceptional circumstances) upon external conditions, is afforded by an appeal to the statistics of deaths resulting from asphyxia in general. Out of 234 cases of death from asphyxia produced in various ways, Maschka found in 48, hyperæmia of the brain, membranes, and sinuses, in 30, a condition rather resembling anæmia, in 156, an ordinary moderate amount of blood.

It is thus evident that a clearly defined condition of the brain and lungs is not to be expected when making post-mortem examinations on cases of death from hanging, nor is it possible from inspection of these organs to determine the relative importance of stoppage of the air- and of the blood-channels as the cause of death in any given case.

Another mode of death from hanging has been described, said to be due to compression of the vagi in the neck by the suspending cord or ligature.

If pressure on the vagi produced any effect on the heart it would be suddenly to arrest it in diastole, there is an abundant evidence, however, apart from physiological experiment, to prove that the heart continues to beat for a considerable time after the stoppage of respiration by hanging. McCausland<sup>2</sup> made some observations on a man executed by hanging, the rate of the pulse being taken every quarter of a minute after suspension. At the third quarter of the first minute it equalled 40 heart-beats per minute, at the last quarter of the fifth minute 152. The radial pulsation became less distinct until the end of the seventh minute, when it could no longer be felt. With the stethoscope the heart-beats could be heard during the eleventh minute at the rate of 120 per minute, at the end of the fourteenth minute it was still heard, but weak and fluttering, at the fifteenth minute it had ceased to beat. Schwab in a similar case found that the heart did not cease to beat until eight minutes after suspension. Maschka<sup>3</sup> observed two similar cases, in one the heart-beats continued for four and the other for five minutes after suspension. Balfour<sup>4</sup> gives three sphygmographic tracings taken from the radial artery of a criminal at intervals of 3½, 5½, and 7½ minutes after suspension, the heart continued to beat for twenty minutes. Misuraca<sup>5</sup> tied both vagi in dogs, and death did not take place until from fourteen hours to even days afterwards. It is not

<sup>1</sup> *Lehrbuch d. ger. Med.*

<sup>2</sup> *Philadelphia Med. and Surg. Reporter*, 1883.

<sup>3</sup> *Handbuch*, Bd. 3.

<sup>4</sup> *Clinical Lectures on Diseases of the Heart*, 1882.

<sup>5</sup> *Rivista Sperimentale*, 1889.



probable, therefore, that pressure on the vagus has to do with the cause of death by hanging.

**Compression of the large vessels of the neck** by arresting the cerebral circulation induces very **rapid loss of consciousness**. The accounts given by those who have been rescued after a sufficiently long suspension to produce complete unconsciousness are mostly to the effect, that after a preliminary feeling of loss of power and ability to make any attempts to move the limbs, which is often preceded by subjective ocular and aural sensations (such as the occurrence of sparks and of a rushing sound), all becomes dark, and consciousness is abolished. The sudden onset of insensibility explains why in accidental and in suicidal hanging no attempts are made by the victim to save himself in many instances the least movement would be enough to avert death. In other modes of committing suicide, voluntary or involuntary endeavours to escape the consequences of the act are not uncommon.

**Judicial hanging** in England is effected by allowing a drop of from six to eight feet, according to the weight of the convict, by which the spine is usually fractured at the level of the second or third cervical vertebræ, compression or laceration of the cord with rapid loss of consciousness results.

**Post-mortem Appearances—External**—Cadaveric rigidity varies as to the time of its appearance. In one case subsequently mentioned it commenced within half an hour after death, in other cases it is delayed. If the body has been long suspended the post-mortem stains are most marked at the lower parts, and there may be punctiform superficial ecchymoses. The face is usually **pale** and the expression tranquil, in some cases the face is swollen and livid. The eyes are half open and usually not prominent, although in some cases they may be, the pupils vary as to dilatation. The tongue may be protruded between the teeth, but this is by no means invariable, it occurs in rather under 50 per cent of cases. Maschka directs attention to a bluish colour of the free border of the lips (which has been noticed by previous observers) as of frequent occurrence—in 98 cases out of 153. According to Adamkiewicz,<sup>1</sup> cyanosis and swelling of face, ears, and lips only occur when the death agony is prolonged, this condition is more likely to be present in plethoric individuals. If the body is cut down at an early period the tumefaction and cyanosis disappear. Punctiform ecchymoses may be present on the face and neck, they are more likely to occur under the conjunctivæ ocular and palpebral and also on the outer surface of the lower eyelids, they are only present in 8 or 10 per cent of cases, therefore their absence is less significant than their presence. Turgescence of the genital organs of both sexes may or may not be present, it is due to hypostatic hyperæmia, and is of no value as a sign of death from hanging. Escape of semen, urine, or fæces is of no diagnostic value, as it results from muscular relaxation, and is met with in many kinds of death both violent and natural. Sometimes saliva flows from the corner of the mouth and may be traced after death, this is not invariably the case, but when observed it indicates suspension during life. The hands may be clenched, but this is at the most an indication of violent death, in suicidal hanging the hands, as a rule, are not clenched unless the act has been associated with more effort than is usually the case, or when the suicide has given himself a long drop.

The most important external appearance is the **mark round the neck** produced by the suspensory medium. Generally speaking, the breadth and depth of the mark depend upon the nature of the ligature used. If it is broad, soft, and yielding, only a superficial mark will be produced, and exceptionally there

<sup>1</sup> *Vierteljahrsschr f ger Med*, Bd 18

may be none at all, if it is thin, hard, and firm, the mark will be deep and narrow. In either case any patterns or irregularity on the surface of the suspensory medium—the strands of a rope, or the texture of a handkerchief, for example—may be reproduced on the skin with which it has been in contact. The mark may entirely surround the neck, but is more frequently limited to the anterior half, and runs between the thyroid cartilage and the hyoid bone, taking an upward direction at the sides immediately behind the ears, where it usually ceases. If a cord with a running noose has been used, and sometimes even with a medium like a pocket-handkerchief, the mark may be continuous round the neck, in the latter case, probably on account of excessive weight of the body or the length of drop. From irregularity of pressure, or from some peculiarity in the nature of the suspensory ligament, the mark may be interrupted in a part where it is usually continuous. Exceptionally the mark may only exist at one side of the neck, the opposite side being free, this arises from the position of the noose, which causes the head to fall sidewise instead of forwards, as is usual. Occasionally the mark runs horizontally round the neck under the thyroid cartilage the original position of the cord being maintained owing to the use of a noose that tightens easily and so keeps its hold and also possibly to unusual prominence of the cartilage preventing the cord from slipping up.

The mark varies, in **colour** and **consistence**, with the nature of the ligature. It may be pale and soft, with slightly reddened or livid borders, or it may be dusky red, or bluish-grey in colour, with margins of a slightly deeper hue, or, finally, it may be yellowish-brown, with darker margins, and of a horny or parchment-like consistence. The first two varieties are the result of soft or yielding ligatures, the last is produced by hard, rough ligatures which injure and partially rub off the epiderm, so that the underlying cutis becomes dry. More than one of these forms may be met with in a single mark—it may be white and soft at one part, brown and hard at another.

**Internal Appearances.**—Hæmorrhage into the subcutaneous tissue in the neighbourhood of the mark produced by the ligature is comparatively rare, Maschka found it only in 10 cases out of 153. The adjacent muscles are not often injured, although Lesser<sup>1</sup> states that in 50 cases of hanging he found injuries to the muscles in 11, mostly in the sterno-mastoid. Fractures of the hyoid bone and thyroid cartilage are of exceptional occurrence, when present, they are mostly due to degenerative changes producing abnormal fragility. Fracture or separation of the vertebræ has only been seen in one or two cases, of which Lesser records one. Rupture of the inner or middle coats of the carotids sometimes happens, Lesser found it in 7 cases out of 50. It mostly occurs in individuals at or past middle life, the 7 cases just mentioned were all above forty years of age, although in none of them was endarteritis deformans present. The remaining internal appearances are common to all modes of death from asphyxia.

The appearance of the **brain** and **lungs** has already been described when discussing the modes of death from hanging. In rather over 50 per cent of cases the amount of blood present in the **brain** and cerebral vessels does not differ from the normal, in the remainder there is sometimes more and sometimes less than normal. Very rarely is there extravasation of blood, although in odd cases small ecchymoses may be present in the dura mater, their presence is significant of the mode of death. The mucous membrane of the **trachea** and **epiglottis** is generally injected. Maschka directs attention to a cyanotic or dark

<sup>1</sup> *Vierteljahrsschr. f. ger. Med.*, 1891.

blue colour of the mucous membrane of the pharynx sharply limited below, which is rarely absent in death from asphyxia and seldom present in other modes of death. The **lungs** may exceptionally contain hæmorrhagic foci in their substance, sometimes they are œdematous, probably in consequence of slow death. Small **sub-pleural extravasations** of blood, known as Tardieu's spots, are very common after death from asphyxia. They are produced by increased blood-pressure (due to asphyxial stimulation of the vaso-motor centre), which leads to rupture of sub-pleural capillaries. In experimenting with dogs 'orn<sup>1</sup> found that, in whatever way the animals were killed, if the blood-pressure was materially increased immediately before death, as demonstrated by a manometer connected with the carotid artery, the lungs at the same time having ceased to move, sub-pleural ecchymoses were invariably produced. Hence the frequency with which they are found after death from asphyxia, the essential factors, elevated blood-pressure with coincident arrest of the movements of the lungs, being usually present. The same factors may occur, however, in other modes of violent death, and may lead to the formation of precisely similar sub-pleural hæmorrhages, thus they have been found in the bodies of individuals suddenly killed by falling from a height, by fire, by CO, and by some other poisons, they may also be found after death caused by certain diseases, such as whooping-cough, purpura, and scurvy. Inasmuch as the occurrence of sub-pleural ecchymoses is not constant their absence is not incompatible with death produced by asphyxia. Ecchymoses sometimes occur in the **heart**, the right half of which is usually filled with dark fluid blood, but its appearance is by no means constant. In the loose cellular tissue about the thoracic aorta and the œsophagus small hæmorrhages are almost constantly present. They are not, however, limited to cases of death from asphyxia, being found after other modes of violent death, and further, very similar appearances may be produced post-mortem in the act of detaching the aorta from the spinal column. Hofmann points out the necessity of washing such hæmorrhages with a gentle stream of water in order to ascertain whether the blood in them is fluid or coagulated, if quite fluid, the hæmorrhages are probably of post-mortem derivation. The mucous membrane of the **stomach** is not infrequently intensely injected, a condition which may be shared by that of the intestines, ecchymoses are occasionally found in both. The **kidneys** are often hyperæmic, sometimes ecchymoses are present. The liver may contain more blood than usual. The **blood** is usually **dark-coloured** and **fluid**.

From the medico-legal standpoint three contingencies have to be considered in cases of death from violence—Was death the result of **Accident**, **Homicide**, or **Suicide**?

**Accidental hanging** is of rare occurrence. With one or two exceptions all the instances recorded have resulted either from playful attempts made by boys to imitate judicial hanging, or, in the case of still younger children, from swinging in the vicinity of a dependent rope, which somehow has caught the child round the neck, and has suspended its body until death took place.

A case of this kind occurred in Norwood in 1895. During the absence of the nurse, a child aged three years got on a stool and was playing at the window, when the stool overturned. The child's neck became entangled in the blind cord, and death took place before assistance was forthcoming. In a still more recent instance a young child amused himself by swinging face downwards, with his chest or stomach balanced in the loop of a clothes line which hung within nine inches off the ground, shortly after, the child was found dead with his neck simply resting on the cord. Maschka records a case of fatal accidental hanging in which

<sup>1</sup> *Archives de Physiol' norm et path*, 1894

a man, slightly under the influence of liquor, slipped off a ladder and was caught in the noose of a rope hanging from the ceiling. Zulch<sup>1</sup> records the analogous fatal case of a boy, aged thirteen, who slipped off a ladder in such a way that a shawl which he wore caught on a hook and suspended him by the neck. Zulch gives another case in which a drunken man fell down some steep steps, and was caught by the neck in the angle which they formed with a post so that he remained suspended until death took place. Kirkhead<sup>2</sup> relates a case which he regards as accidental rather than suicidal, in which the mode of suspension was peculiar. A young man, eighteen years of age, who was a prisoner, was found dead in his cell suspended by one of his braces round the neck, his feet touching the floor, the braces were made of iron and lined with coarse calico. The deceased, with his back to the wall, had evidently stood on some hot water pipes about 6 inches from the floor, placing the brace under the lower jaw he had passed the ends, without tying them, through a chain-loop which depended from the window of the cell, he had then either stepped or slipped off the pipe, when the sides of the chain loop came together sufficiently close as to grip the brace and hold it fast, with the body suspended by it. The brace was not knotted nor fastened to the chain in any way, and it appeared highly improbable that it could in this manner sustain the weight of the body, subsequent trial showed that the grip of the chain was sufficient to prevent the rough surfaced material from slipping under the strain placed upon it. The incompleteness of the preparations, with other circumstances, made it probable that the prisoner contemplated only a feigned attempt at suicide in order that he might be placed in a ward with company. Quick onset of cadaveric rigidity was observed in this case, the boy was seen alive at 2.20, the body was found at 2.30, and at 3 o'clock cadaveric rigidity had commenced in the neck.

A most exceptional case of accidental hanging was the occasion of a trial for man slaughter, *Reg. v. Montague* (Com. Oyer and Term. Dub., 1892). The prisoner was accused of having caused the death of her daughter, a child three years old. As a punishment, she tied the child's arms above the elbows with a stocking, which was then passed round the body so as to pinion them to the sides. At the back one end of a cord was attached to the stocking, the other end being fastened—5 feet 8 inches from the ground, and, consequently, about 2½ feet above the child's head—to a ring in the wall of a dark closet. The child was left three hours, and when the mother went to liberate her she found her dead and suspended by the cord in such a way that when it was loosened from the ring the body fell forward. There was a mark produced by pressure on the neck at the lower part of the windpipe. Either the stocking slipped up to the throat, or the cord got partially round the front of the neck, the result being death from hanging. The prisoner was found guilty, and was sentenced to twelve months' imprisonment.

**Homicidal hanging** is also very infrequent. It would be exceedingly difficult to murder a man of average strength by hanging unless he was deprived of the power of resistance. In the case of the very old or very young it may be accomplished single-handed, or even with a robust man if he was taken unawares. In the greatest number of cases, however—not the result of accident or suicide—in which bodies are found hanging death has been caused in some other way, and the body hung in order to simulate suicide. This leads to a consideration of the signs of death from hanging with regard to the possibility of distinguishing between **suspension during life and after death.**

First, as to the punctiform ecchymoses that may be found on the lower parts of the body. Two antagonistic views are held as to their causation, one is that they are produced after death by the weight of the blood in the body rupturing the coats of some of the smaller vessels, the other is that they are formed during life from the increased blood-tension occasioned by the asphyxia. Observations are forthcoming which serve to support both views. Lesser<sup>3</sup> hanged recently-dead bodies for twenty-four hours, and although the lower parts were as hyperæmic as they well could be, not a trace of extravasation was found, he also found extravasations in the non-dependent parts of bodies in which death had resulted from asphyxia. Strassmann<sup>4</sup> relates a case of suicidal hanging in which the body was found in a kneeling posture, the post-

<sup>1</sup> *Zeitschr. f. Med. Beamte*, 1894.

<sup>2</sup> *The Lancet*, 1885.

<sup>3</sup> *Vierteljahrsschr. f. ger. Med.*, 1884.

<sup>4</sup> *Ibid.*, 1888.

mortem stains were limited to the thighs, where ecchymoses were found in abundance, the legs below the knees were pale and free from ecchymoses. Hofmann<sup>1</sup> believes that the extravasations are formed *intra-vitam*, but developed after death when the body is left suspended. Linn<sup>2</sup> regards the ecchymoses solely as indicative of prolonged suspension after death. It may be accepted that ecchymoses on the lower part of the body are of no value as indications of suspension during life.

A mark round the neck affords no *proof* that death was caused by hanging. Casper regarded the mark as purely a post-mortem phenomenon, probably a too sweeping assertion, still it is perfectly true that a cord-mark may be produced after death that cannot be distinguished from one produced during life, this applies not only to a period immediately after death, but also to an interval of several hours after. Punctiform extravasations of blood are occasionally found when the cord-mark is cut into, but they cannot be relied on to indicate suspension during life, as they have been found in the bodies of those suspended after death, for the same reason the injected vessels along the borders of the marks are of no diagnostic value. Deeper-seated injuries in the neck, such as fracture of the cartilages of hyoid bone, or rupture of the muscles, have also been found after post-mortem hanging. Rents in the inner coats of the carotids when accompanied by extravasation of blood within the walls of the arteries, are indicative of suspension during life, they occur very rarely, however, and are easily caused accidentally when making the post-mortem examination. It is to be remembered, on the other hand, that death may result from hanging and yet, for the reasons previously given, no trace of a mark round the neck may be visible. The presence on the chin of saliva which has flowed from the mouth points to suspension during life, but it is not a constant sign, nor one on which alone a decided opinion can be expressed. The remaining internal signs are at the most indications of death from asphyxia – they cannot be utilised as evidence of death from hanging.

The result of this inquiry as to the value of the signs of death by hanging in determining whether a body was suspended before or after death is, that they are *not* sufficiently conclusive as to justify the expression of an absolute opinion.

Other circumstances should be taken into consideration in attempting to solve the question. The body should be carefully examined for signs of struggling or of injuries other than those due to the assumed mode of death, the possibility of poisoning must not be overlooked. The absence of any such signs is more significant than their presence. It is not an unfrequent occurrence for a person who has determined to commit suicide to make the attempt in more than one way, such a person may ineffectually cut his throat or take poison, or in some other way injure himself, and finding that death is long in coming, may subsequently effect his purpose by hanging. If the injuries are of such a nature that they could not be self-inflicted, the fact of the body being suspended affords presumptive proof of homicide. The possibility of accidental post-mortem injuries caused by the body falling to the ground when cut down, or, if examined at a distance from the place where it was found, by rough handling during transport, should not be lost sight of.

Illustrations of self-inflicted wounds found on bodies in cases of suicidal hanging which might at first sight give rise to suspicion of homicide are given by Maschka. In one case, several incised wounds were found on the inner side of the left forearm, one of which divided the radial artery. In another case,

<sup>1</sup> *Med. ch. L. Ges. Med.*

a suicide discharged a firearm into his mouth, fracturing the hard palate and lacerating the soft parts and then hanged himself. In a third, an old woman, 60 years of age hanged herself, and was cut down by her husband, on return to consciousness she seized a knife and cut her throat. At the necropsy an ordinary cut-throat wound was found above the thyroid cartilage and a cord mark as well.

In some cases the balance of probability would be against that which actually occurred. Lman<sup>1</sup> records the case of a woman who inflicted on herself two wounds which penetrated the pericardium and afterwards hanged herself also that of a man who in consequence of being shot in the back by another person hanged himself. In such cases it would be impossible to determine the question of suicide or homicide, from the post-mortem appearances alone. Compare them with some cases of homicide in which the dead body was subsequently hanged, and the necessity for great caution is apparent. Deveaux relates the case of a woman who was found hanged in a barn, on careful examination a small round wound was found under the left breast which transfixed the heart the case being one of homicide. In another case, reported by Vrolik, a sailor was stabbed through the heart by a woman in a brothel, the body was afterwards washed, clad in a clean shirt, and hanged to make it appear that the death was suicidal.

Homicidal hanging is almost exclusively confined to young children, in adults, strangulation and suffocation—the analogous modes of death—are much more easy of accomplishment. One case is recorded by Ogston in which a woman tied a ligature round the neck of her husband while he was asleep and then pulled him up with it. Hofmann<sup>2</sup> relates the case of a man who, by hanging, killed five of his children, aged 8 months, 2, 6, 8 and 9 years respectively, and finally hanged himself. In another case a man hanged his two children (girls), aged 6 and 13 years, probably whilst they were asleep.

In 1888 an exceptional and dramatic mode of homicidal hanging was put into execution in Paris by a man named Eyraud with the aid of a female accomplice named Bompard. The girl formed an illicit connection with one Gouffe, who was the selected victim. In the alcove of a room where an interview was to take place between Gouffe and Bompard, Eyraud fixed a compound pulley, over which a rope furnished with a strong hook was passed, the apparatus and the alcove were concealed by a curtain behind which Eyraud was placed, in front of the alcove was a sofa. Whilst Gouffe was sitting on the sofa the girl placed herself on his knee and in a playful manner adjusted the noose of a silk cord round his neck, and then passed the free end of the cord (which was furnished with an eye or small loop) to her accomplice, who slipped the eye over the hook depending from the pulley, and pulled the rope to which it was attached until Gouffe was drawn up from the sitting posture and suspended by the neck, the rope was then made fast to the sofa so as to keep the body suspended. After robbing the corpse—which was the object of the murder—it was put into a box and taken to a distance and then left. The body was discovered fourteen days after, and on examination it was found that both cornua of the hyoid bone were fractured. The culprits were ultimately arrested, and from the confession of Bompard the actual mode in which the deed was perpetrated became known. The case is remarkable in itself, and further has an adventitious interest, because the question of hypnotic suggestion was raised in favour of the female prisoner, it was urged that she was subject to the will of her accomplice, and was not personally responsible for the part she took in the transaction. A full account of this case by A. Lacassagne is contained in the *Arch. de l'Anthropologie Crim.*, 1890.

**Suicidal Hanging.** Hanging is one of the commonest modes of committing suicide, judicial hanging apart, suicide is accountable for more than 99 per cent of the total deaths by hanging. In 1919, the deaths of 649 men and

<sup>1</sup> *Handbuch d. ger. Med.*

<sup>2</sup> *Lehrbuch d. ger. Med.*

185 women were attributed to suicide by hanging and strangulation in England and Wales. The returns of the Registrar-General do not differentiate between these two causes of death, but strangulation would only have accounted for a small proportion of the total. Six of these deaths (four male and two female) were between the ages of ten and fifteen years. The reason why this mode of self-destruction is relatively so frequently adopted is that the means for carrying it out are to be found in every house, and but little skill or effort is required in its execution, and further, the act of placing the head through a noose and allowing the body to depend from it partakes of a more passive character than that of using a knife to cut the throat or a revolver to blow out the brains.

If the body of an adult of ordinary physical development is found hanging without any sign of injury or of struggling, and the general appearances are consistent with death from hanging, the probability of its being a case of suicide is very great. From what has been previously said, the presence of wounds or of poison does not preclude suicide, but in such cases the contingency of homicide would be greater. In addition to making a thorough examination of the surface of the body for marks due to scratches, bruises, or blows, the clothing and immediate surroundings should be investigated. A few scratches on the neck are not necessarily indicative of homicide, occasionally they are inflicted by the victim himself involuntarily clutching at the noose. Care must be taken not to mistake signs of antecedent injuries for those caused at the time of death.

No conclusion can be arrived at as to the relative probability of suicide or homicide from the position in which the body is found. It is not necessary that the feet should cease to rest on the ground, sufficient pressure on the neck to cause death may be produced even in the sitting posture. Grant<sup>1</sup> relates how a man, aged 48 years, hanged himself by tying the ends of a common cotton pocket-handkerchief together and suspending it from the handle of his bedroom door, which was only 2 feet 9 inches from the floor, he then sat down on the floor with his back to the door, put his head through the loop and bent forwards. This was proved by flattening of the nates—cadaveric rigidity having set in before the body was discovered—and by the post-mortem staining on the lower part of the back and legs. The mark on the neck was 1½ inches broad and extended from below the chin obliquely upwards and backwards over the occiput, the texture of the cloth being plainly impressed on the skin. Many cases of a more or less similar nature are recorded. Nobiling<sup>2</sup> reports the case of a man, aged 24, who hanged himself with a pocket-handkerchief attached to the latch of a door, which was 3 feet 7 inches from the floor. He adopted the kneeling posture and was found dead with the knees bent and the toes touching the floor. The handkerchief was folded broadly and left no mark behind.

In some cases the attitude is such that death can scarcely be said to be due to hanging, inasmuch as there is little or no suspension, such cases more nearly approach strangulation. Hurry<sup>3</sup> records a case of this kind, of which the accompanying figure is an illustration.

A woman, 77 years of age, was found dead in the posture depicted. The arms were extended alongside the trunk, and the pronated hands touched the floor on their dorsal aspect, the legs and feet were also extended, the night-dress and chemise covered the body and showed no signs of disorder. The

<sup>1</sup> *The Lancet*, 1889

<sup>2</sup> *Aerztliches Intelligenzblatt f. Baiern*, 1884

<sup>3</sup> *Annales d'Hygiène*, 1881

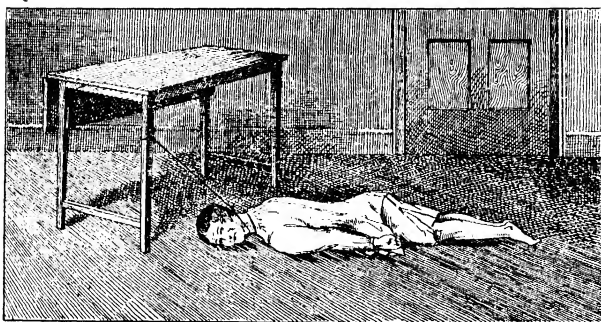


Fig. 16.

cord by which she hanged herself was nearly 40 inches long, and was attached about 17 inches from the floor to the leg of a heavy table. The head half-rotated

to the left rested with the right malar prominence on the floor, about 12 inches from the table foot. The woman was subject to delusions, and had committed suicide in this exceptional manner. In Tardieu's *Etude sur la Pendaison* a number of somewhat similar instances of hanging by incomplete suspension are illustrated.



Fig. 17.

The diagnosis between suicide and homicide is occasionally embarrassed by the hands or the legs of the deceased being found tied together; in such cases the way in which the limbs are secured, the position and the kind of knot, and the firmness with which it is tied, are to be carefully noted. In some instances there is little difficulty in determining that the ligature round the limbs was tied by the victim himself; in other undoubted cases of suicide the position or the character of the knot is such as to make it appear almost impossible that it was self-tied. A striking illustration of the latter kind is given by Filippi,



Severi, and Montalti<sup>1</sup> The nude dead body of a man hanging by the neck was found in his office, the right foot touched the floor, the left was slightly lifted from it by the flexion of the knee The wrists, tied together, were behind the back in such a way that the knot of the cord was in front of them and rested on the back of the body It was exceedingly improbable that the knot could have been tied behind the back by the individual himself, and as every other indication pointed to suicide, the question arose could he have tied them in front, and then passed his legs between the arms? Professor Filippi, to whom I am indebted for a copy of a photograph taken at the time, informs me that subsequently the question was solved by a young acrobat, who with considerable difficulty, succeeded in passing his legs between his arms, with the wrists tied together in front

**The After-effects of Threatened Death from Hanging** When the body is cut down before life is extinct, efforts at resuscitation may be so far successful as to re-establish respiration, it is to be noted, however, that in some cases return of automatic respiration is not followed by ultimate recovery the surface of the body remains cold the circulation feeble, and insensibility persists, death taking place some hours subsequently When recovery takes place certain neuroses or psychoses occasionally manifest themselves Terrien<sup>2</sup> met with one case, in which epileptiform convulsions with tetanic spasm drawing the head to one side occurred the patient remained unconscious for several days, and on regaining consciousness was found to be amnesic In a second case there were epileptiform convulsions with opisthotonos, and on return to consciousness the patient for some time made movements as though walking Petina records the case of a man who remained unconscious for twenty-four hours after attempted suicide by strangulation, during which time he had violent clonic convulsions of the whole body, but especially of the right side of the face and of the left arm, followed by a general condition of muscular rigidity When he became conscious, the speech was stammering but deglutition was not difficult, there was paralysis of the right facial nerve and of the left side of the body, with diminished sensibility on the right side of the face and on the left side of the body, except on the front of the thorax, when there was some hyperæsthesia The paralysis gradually improved, but ataxia subsequently occurred on the non-paralysed side, the inco-ordination spreading from the upper to the lower extremities, and then passing over to the paralysed arm This was probably a case of hæmorrhage into the pons Wagner<sup>1</sup> records the occurrence of acute dementia as a result of threatened death from hanging

## STRANGULATION.

When death results from asphyxia caused by constriction of the neck by means of some form of ligature without suspension of the body, it is said to be due to strangulation The mode of death is the same as that which occurs in hanging

**Post-mortem Appearances External**—The only appearance that needs special consideration is the mark produced by the ligature round the neck As a rule, the mark produced by strangulation differs from that produced by hanging in taking a more horizontal direction and in more completely encircling the neck, since the direction of the constrictive force does not necessarily tend to draw the ligature above the thyroid cartilage as in hanging, the mark is not infrequently on the level of or below the cartilage

Exceptions occur when the victim is thrown to the ground and the ligature is pulled upwards and backwards, the direction taken by the mark then resembles that due to hanging On the other hand, a horizontal mark may be produced by hanging when the position of the noose is different from that which is usual—e.g., the knot being under the chin The nature of the ligature

<sup>1</sup> *Manuale di Med. Legale*, 1889

<sup>2</sup> *Progrès Médical*, 1887

<sup>1</sup> *Prag. med. Wochenschr.*, 1880

<sup>4</sup> *Jahrbucher f. Psychiat.*, 1889

influences the character of the resulting mark if the ligature exercises unequal pressure as it surrounds the neck, the mark will be irregularly formed and may be interrupted in its course. Death has resulted from strangulation without the occurrence of any mark in such cases it is probable that the ligature was of a yielding nature, was not applied with excessive force, and did not remain long round the neck.

The observations already made as to the **absence** of differential signs between marks round the neck produced by suspension of the body before death and after apply equally to marks due to strangulation.

**Internal appearances** resemble those described as due to death from hanging.

**Accidental strangulation** was responsible, in 1919, for sixteen deaths of males and four deaths of females in England and Wales. The circumstances of such deaths vary. Children have been strangled by playing with blind-cords, infants by mufflers tied too tightly round the neck, and epileptics, drunkards, and other helpless persons getting into positions which have led to compression of the neck. Taylor and Stevenson record the case of a girl who was employed in carrying fish in a basket supported on her back by a leathern strap passing above the shoulders round the front of her neck. She was found dead sitting on a stone wall, off which the basket probably slipped while she was resting, causing the strap to compress firmly the windpipe. Gordon Smith<sup>1</sup> gives a similar case of a boy who was in the habit of going about with a cord round his neck, to which a weight was attached, he was found dead in a chair, the weight having changed its position and drawn the cord tight round the front of the neck. Lesser<sup>2</sup> records the case of an epileptic, who was found dead with his feet on the bed and his face and breast on the floor, he was clad in his shirt only, which was gripped between his body and the lower part of the bed so that the collar was tightly drawn round the neck. On the lower part of the neck there was a shallow mark rather over half an inch broad, not depressed but sharply defined from the hæmorrhages and post-mortem staining in the surrounding skin on the face and lower part of the neck were punctiform ecchymoses. As Lesser admits, this is a doubtful case, as although there were indications of asphyxia, it is by no means clear that strangulation was the cause of it, the mouth and chest on the floor (which was covered with a carpet), and the epileptic seizure, are sufficient to account for the asphyxia and the ecchymoses, the mark round the neck might be due to pressure from the collar rendering the underlying skin bloodless without arresting breathing. At the Manchester summer assize 1895 a man was indicted for having caused the death of a boy aged thirteen. The prisoner caught the boy trespassing in a garden and placed him in a stable, fastening a chain, by means of a padlock, loosely round his neck, the other end of the chain being attached to a post, a short time after the boy, in a sitting posture, was found dead. The post-mortem examination showed that death took place from suffocation. The prisoner was discharged. In 1900, near Manchester, a child which was left by her mother in bed, was soon after found dead on the floor, strangled with the strings of her nightdress, which by her struggles she had drawn tightly round the throat.

**Homicidal Strangulation.**—The relative frequency of homicidal and suicidal strangulation is the converse of that which occurs when hanging is the cause of death. From 1885 to 1894, in England and Wales, 147 persons were murdered by strangulation, of these, 121 were under the age of one month, 5 under the age of one year, and 5 under the age of five years. In the following decade, 1895 to 1904, 157 persons were murdered by strangulation, of these, 128 were

<sup>1</sup> *Principles of Forensic Med.*, 1827.

<sup>2</sup> *Atlas der ger. Med.* (Zweite Abtheilung), 1890.

under the age of one month, 5 under one year, and 4 under five years. During the seven years 1905 to 1911 the number of such deaths was 123, of which 80 were of children below three months. On account of the rapidity with which insensibility supervenes when the throat is constricted, as is the case in strangulation even a strong man, if taken unawares, is unable to defend himself or cry for help. It is much easier to strangle a man than to hang him, because the constrictive force can be brought into play the moment the ligature is placed round the neck, whereas in homicidal hanging the victim has time to resist, because the constrictive force does not come into play until the body is lifted off the ground, and in the meantime he is put on the alert by feeling the cord round the neck. A case described by Lesser<sup>1</sup> shows how easily homicidal strangulation may be effected even when the victim is a strong man. A man twenty-seven years of age, in complete possession of his bodily and mental powers, had a cord thrown round his neck from behind whilst sitting on a chair, the ends were drawn together, carried twice more round, tied once in a single knot in front, and a second time in a double knot at the angle of the right lower jaw, the victim fell senseless without making any attempt at resistance, the deed being perpetrated by one man. An adult has been strangled in a room divided only by a wooden partition from another in which were persons who heard no suspicious sounds.

In accordance with the general principle that *a murderer uses more violence* than is necessary to effect his purpose, the distinctive feature of homicidal strangulation is greater local injury than is found in strangulation from accident or suicide. A diagnosis is chiefly required between homicide and suicide, accidental strangulation is rarely alleged in defence in cases of suspected homicide, as the surroundings would probably determine the question off-hand. Mere extravasations of blood under the mark in the neck, though pointing to, are not indicative of homicide nor are external injuries, such as scratches, excoriations of the skin, and slight bruises of more significance, they are suspicious indications, but nothing further. On the other hand, fractures of the thyroid or cricoid cartilages, or of the hyoid bone, are distinctly indicative of homicide, as they are rarely if ever, found in cases of suicidal strangulation, unfortunately for diagnostic purposes such injuries are frequently absent in homicidal strangulation. Bleeding from the ears has been caused by strangulation accompanied with excessive violence, but it is of exceptionally rare occurrence. It is not necessary that the tympanum should be ruptured, although this has happened, the blood may be derived from the meatus, external to the tympanum, from bursting of some of the small superficial vessels in the same way that subcutaneous extravasations are formed, the effused blood obtaining exit through slight rents in the skin. Bleeding from the ears without rupture of the tympanum has, however, been observed in suicidal strangulation. Protrusion of the tongue against or between the teeth, and of the eyeballs, is more likely to occur in violent strangulation than in hanging, but is not a constant sign. Turgescence of the face with a cyanotic or dusky red hue may also occur, but, as with protrusion of the eyeballs, if present at the moment of death, it may disappear on removal of the ligature, even after strangulation with violence the face may be pallid. Blood has been observed to escape from both mouth and nostrils after homicidal strangulation, in such cases it is probable that the violence of the act is not the sole cause, but that extreme efforts at self-defence on the part of the victim unduly distend the blood-vessels of the head before the ligature is sufficiently tightened as to arrest completely the

<sup>1</sup> *Atlas der ger Med (Zweite Abtheilung)*, 1890

blood current, in suicidal strangulation, bleeding from the nostrils has occurred from the violence of the efforts made by the suicide to tighten the ligature to the utmost.

**Signs of general violence** are to be sought for as indicative of homicide. As previously stated, scratches or excoriations in the neighbourhood of the strangulation mark are suspicious but do not constitute proofs of homicidal violence—they may have been produced by the ligature before it arrived at its final position or by the suicide involuntarily plucking at it especially if the finger-nail are long. The rest of the body should be carefully examined for wounds and bruises—the back of the shoulders, for example, may show traces of counter-pressure employed to steady the victim whilst the ligature was being tightened. When discussing the subject of supposed suicidal hanging, it was mentioned that death may be caused by various kinds of violence—the body being subsequently suspended by the neck to simulate suicide by hanging. In such cases, when death has been caused by strangulation there would probably be two marks on the neck—this, however, has occurred in suicidal hanging from the neck slipping further through the noose after a limited period of suspension.

Homicidal strangulation occupies a prominent position among the modes of violent death that from time to time are **feigned** by an individual who wishes either to excite compassion—to account for the loss of a sum of money entrusted to his charge which he has embezzled, but which he alleges was stolen from him after he was rendered unconscious—to revenge himself on some one against whom he has a grudge by accusing him of a homicidal act—or in the case of hysterical women without any obvious reason whatever. All cases of alleged attempted homicidal strangulation, in which the victim has sustained little external injury should be approached with a certain amount of scepticism on the part of the medical man which, however, need not be displayed in the first instance—the mental attitude should be that of a man open to conviction but who requires convincing. Many of these cases of imputed homicidal violence would scarcely be heard of were it not that the medical man who first sees the case does not approach it in a sufficiently judicial frame of mind—he accepts the statements of others who first discovered the supposed victim without sufficiently weighing their probability. Tardieu relates several instances of this description—in one, a man, stated to be in an almost lifeless condition, was found in a cellar with the hands and feet tied, and a cord three or four times round the neck, but not tied—in three hours he was himself again, except that he could not speak. According to the man's own statement subsequently made, he had been eleven hours in the condition as found, and although the whole affair was a palpable fraud, still medical evidence was forthcoming in favour of the account given, which was that the man's master had attempted to strangle him. Another case was that of a girl who asserted that she was the victim of a political conspiracy, and that a man had attempted to strangle her, and at the same time had stabbed her with a dagger, her dress was cut, but not her body. She was dumb when discovered after the alleged assault, but when Tardieu told her she would speak in a minute, she did so and confessed the imposture.

**Suicidal Strangulation**—Although strangulation is an exceptional mode of committing suicide, instances are not so rare as formerly was assumed, self-strangulation was regarded as next to impossible, except some contrivance of the nature of a tourniquet was adopted such as passing a short stick between the cord and the neck, and twisting it round until asphyxia was produced. Numerous cases have proved, however, that the necessary constrictive force

can be brought to bear by simply encircling the neck once or twice with a ligature and then tying it, or even without tying the way in which the ligature is applied, together with the position and the formation of the knot, are of great importance in differentiating between suicidal and homicidal strangulation. Maschka considers that when the ligature goes several times round the neck, suicide is indicated rather than homicide, because quick and powerful tightening of it by a murderer would be more easily accomplished with a single turn than if there were several. On the other hand, there is nothing to prevent a murderer from using a single turn first, and, when unconsciousness has been produced, twisting the cord once or twice more round the neck, and tying it to make sure that the victim shall not recover. In Lesser's case of homicidal strangulation, previously mentioned, the cord went several times round. The position of the knot varies—it is most commonly found in front or to one side, usually the left, exceptionally it has been found on the back of the neck, the right- or left-handedness of the suicide may affect the position of the knot. More than one knot, especially if one is separated from the other by an extra turn of the cord round the neck, points to homicide and the more perfect the formation of the knot, the less likely is the case to be one of suicide. Anything peculiar about the construction of the knot must be noted—sometimes an indication is thus obtained as to the trade of the person who made it—sailors, for example, use knots of a kind different from those used by landmen, the material of which the ligature consists may also afford a clue.

The **degree of violence** inflicted by the ligature is a valuable indication. Fractures of the thyroid or cricoid cartilages or of the hyoid bone may almost be taken to exclude suicide, the like may be said of rupture of the muscles and of the inner coats of the carotids. Slight extravasations into the substance of the muscles underlying or near the cord-mark may occur in suicidal strangulation, but they are then limited in extent being for the most part of the punctiform type. When the subject of homicidal strangulation was discussed, attention was drawn to the possibility of after suspension of the body in order to simulate suicide, and the consequent formation of a double mark round the neck, a like occurrence may be met with when a primary and unsuccessful attempt at suicide by strangulation has been followed by self-suspension. A careful comparison of the two marks would probably enable an opinion to be formed as to the real mode of death—if by strangulation, the lower and more horizontal mark would be characterised by signs of greater violence than the mark due to suspension, if by hanging, the converse would hold good.

Signs of **general violence** would be **absent**, except such as might be produced in applying the ligature—scratches and slight abrasions about the neck. The possibility of unsuccessful attempts to put an end to life in other ways before adopting strangulation must not be lost sight of, when the body is examined before it has been disturbed, the state of the clothing and of the surrounding objects would show whether there had been any struggling or not. It is to be remembered, however, that a murderer might obliterate such indications, and also that suicides have been known to be affected with a kind of fury before putting an end to themselves, and have thrown all the surroundings into disorder.

The following cases are illustrative of the manner in which suicidal strangulation has been effected. In a case reported by Filippi,<sup>1</sup> an insane man strangled himself by simply surrounding his neck twice with a strip of woollen material and tying it in front. Francis<sup>2</sup> records the case of a man, also a lunatic, who

<sup>1</sup> *Rivista Sperimentale di fren. e med. legale*, 1879

<sup>2</sup> *Med. Times and Gazette*, 1876

strangled himself by twisting some string round his neck, tying the ends to his wrists, and then extending his arms to the utmost, he was found dead, having fallen from his original kneeling posture on to one side. In a case related by Ogston,<sup>1</sup> the body of an old man was found in a wood with a neckcloth wound more than once round the neck, a walking-stick had been passed through the loop of the neckcloth and twisted so as to tighten it. From the obliquity of the outer fold of the neckcloth, Ogston at first thought that the body had been dragged along the ground with the stick, on careful examination he concluded that the oblique direction of the neckcloth had been produced by a final conclusive struggle, the stick, entangled in a clump of young trees, had held the ligature fast, whilst the movements of the body had drawn it into the position in which it was found.

### STRANGULATION BY THROTTLING.

When strangulation is effected with the unaided hands, the act is called throttling.

The experiments made on the dead body by Langreuter (see section on *Hanging*) illustrate the facility with which the air-way may be occluded by external pressure. It was found that if the thumb and forefinger were placed externally on the two sides of the thyroid cartilage respectively, the least pressure was sufficient to close the aperture between the vocal cords, stronger pressure made the cords overlap each other. When strong pressure was made between the larynx and the hyoid bone, the air-passage was stopped, principally through approximation of the ary-epiglottic ligaments.

The only points to which attention need be specially directed in relation to throttling are the external and internal appearances produced by the pressure of the fingers on the throat. The general external and internal appearances are those common to other forms of strangulation.

**Position of the Marks.**—They are usually found on both sides of the front of the throat directly below the lower jaw, sometimes they are far back, at the angle of the lower jaw. They may be sufficiently detached as to indicate the exact spot occupied by each finger, but not unfrequently they are too closely clustered together to be separately identified. They are distinguishable from an interrupted mark of a cord by not being on the same level. The marks may correspond more or less with the shape of the finger tips, but are often irregular in outline, from infiltration of extravasated blood round about. The impressions of the finger nails may be present as crescentic indents, and if there has been much struggling the skin of the throat will be scratched and excoriated. There may be only one mark on one side and several on the other, respectively corresponding to the thumb and fingers, if the assailant was right-handed, the thumb mark will be on the right side of the neck, if the left hand was used, it will be on the left side. The appearance of the marks, if examined not long after death, is that of bruising, more or less blood escapes beneath the skin and produces the characteristic discoloration. If some time elapses before the examination is made, the marks may present the dry, horny, or parchment-like look and feel, described in one form of the cord-mark produced by hanging, this is due to injury sustained by the epiderm from rough contact with the fingers, which allows evaporation to take place from the denuded surface of the cutis.

<sup>1</sup> *Lectures on Med Jurisp*, 1878

**General signs** of violence will probably be visible in some parts of the body, the attack is often made on a victim in the erect posture, and in falling injuries to the head are easily sustained

**On section** a considerable amount of extravasated blood is usually found in the underlying soft tissues, fractures of the thyroid and cricoid cartilages and of the hyoid bone are frequently met with the carotids escape injury. Lacerations of the inner surface of the cheeks from violent pressure against the teeth may occur

It is usually asserted that throttling is necessarily and invariably a homicidal act. Accident as a mode of production may be ignored, and it has been held to be an impossibility for any one to commit suicide by compressing the throat with the unaided fingers. Whilst allowing that for all practical purposes **throttling** may be regarded solely from the **homicidal** standpoint, still, one solitary case of suicidal throttling is recorded by Binner<sup>1</sup>. A woman, aged forty, suffering from melancholia, who had previously made several attempts to commit suicide, was found dead, crouched in her bed with both hands compressing her throat, the elbows were supported on the knees, and the back leaned against the wall, there were marks of her finger nails on both sides of the throat. Death resulted from compression of the throat by the fingers.

A case of homicidal throttling was tried at the Liverpool winter assize in 1884. A man and his wife had been in the house a short time together when the man went out and sought assistance, "because his wife had fallen into the fireplace and he feared she was dead." She was found apparently dead, sitting in a chair, with the head supported by the wall, marks of blood were on the table, the wall, and the floor of the room. On examination of the body a bruise was found immediately beneath the lobule of the left ear, and another three-quarters of an inch below the right ear, superficial extravasation was evident in both, together with a second and deeper extravasation half an inch below the surface of the right-hand bruise. Other bruises were found over each eyebrow, at the back of the right wrist, over the knuckle of the left little finger, at the inner part of the left elbow, and at each angle of the mouth. The tongue and gums were also injured, and one of the teeth in the upper jaw was partly loosened, the deeper structures of the throat were not injured. The internal signs were those of death from asphyxia, the brain and membranes were intensely congested. The man was found guilty.<sup>2</sup>

## SUFFOCATION.

When air is prevented from entering the lungs by other means than by external constriction of the throat, or by submersion in water, if death results, it is said to be due to suffocation.

The modes of death from asphyxia, which have previously been considered, are necessarily solely due to external influences brought to bear either accidentally or intentionally, from these suffocation differs, inasmuch as it may result from pathological causes apart from external agency. It is not necessary to enumerate all the morbid conditions which may lead to death from suffocation, a few will serve as illustrations. —In children, spasm of the glottis or accumulation of exudative products in the air-passages, in adults, oedema of the glottis, paralysis of the vocal cords, bursting of a pharyngeal abscess or of a thoracic aneurism into the trachea may suddenly cause death from suffocation. Nothing

<sup>1</sup> *Zeitschr. f. med. Beamte*, 1888

<sup>2</sup> *Med. Chron.*, 1885

more need be said with regard to these and like conditions, except that the possible occurrence of those which leave no evidence after death must not be forgotten when searching for the cause of death in doubtful cases.

Suffocation may be produced from external causes by (a) the introduction of **foreign bodies** into the **air-passages**, (b) forcible **compression** of the **chest**, (c) **covering** over the **mouth and nose** (*smothering*).

(a) The introduction of **foreign bodies** into the **air-passages** from **accidental** causes may occur either in the case of adults or of children. The foreign substance is usually some kind of food, either solid or pultaceous, it may be introduced directly, as in the act of deglutition, or it may be regurgitated from the stomach, and find its way into the air-passages instead of being ejected from the mouth. When food enters the larynx, it is frequently the result of an involuntary movement of inspiration induced by sudden surprise or by a fit of laughter during the act of deglutition. A man who is intoxicated may vomit, and, through diminished reflex activity, no effort may be made to prevent the vomited matter from being drawn into the larynx. This may happen apart from intoxication. Coutagne<sup>1</sup> relates the case of a woman aged forty-four, who, feeling unwell, went into a chemist's shop and vomited on entering, she became cyanosed and straightway died. An autopsy revealed that the stomach was greatly dilated, and that it contained more than two quarts of semi-fluid food, some of which had found its way into the œsophagus, pharynx, mouth, and nasal cavities. From the glottis to the bifurcation of the trachea the passage was free, the bronchi down to the smaller branches were completely obstructed. No sub-pleural ecchymoses were present, but the lungs were very large, and on section, serum mixed with froth escaped so abundantly as to resemble the appearances produced by the pulmonary œdema due to drowning. It must be borne in mind that it is quite possible for some of the more fluid contents of the stomach to be accidentally forced into the upper air-passages during removal of a body after death, their presence, however, in the finer bronchi, as in the above case, would afford evidence of vital inspiration. Children are easily suffocated in this way. Mitchell<sup>2</sup> relates the case of a girl aged five, who, after dinner, ate an orange, and feeling sick was put to bed. Shortly after she was found dead, some vomited matter being on the bed-clothes, the child's mother had been in the room all the time without hearing anything amiss. On opening the larynx a piece of orange pulp was discovered impacted in the rima glottidis. In some cases death occurs with exceeding rapidity, especially if the individual is advanced in years, and on post-mortem examination the signs of death from asphyxia are found to be entirely absent, there is neither cyanosis during life, nor excess of blood in lungs, brain, or right heart after death. It is probable, in cases of this description, that death results from syncope. Perrin<sup>3</sup> directs attention to the absence of the post-mortem signs of death from asphyxia under conditions which are likely to produce them, and gives illustrative cases. An invalid, 62 years of age, in the midst of a meal suddenly fell on the knees of his neighbour quite dead. On examination, the epiglottis was erect and rigid, and a mass of food as large as a hazel-nut was found in the glosso-epiglottic fossa, at the fifth ring, the lumen of the trachea was obstructed by a cylindrical mass nearly 2½ inches long which reached down to the point of bifurcation. The lungs were normal and free from ecchymoses, the left side of the heart was empty, and the right side only contained a few spoonfuls of partially fluid blood. In another case a man 68 years of age while leaving a café suddenly

<sup>1</sup> *Annales d'Hygiène*, 1893

<sup>2</sup> *Brit Med Journ*, 1895

<sup>3</sup> *Poulet on Foreign Bodies in Surg Practice*, 1881



fell as though struck by lightning. At the necropsy, a mass composed of pancake was found to fill the posterior part of the pharynx, and to extend to the glottis, the epiglottis was raised.

It is not necessary that the air-passages should be blocked up with the foreign substance in order to induce suffocation, the introduction of a small extraneous body in such a way that it lodges on or between the vocal cords may be sufficient to excite spasmodic closure of the glottis, and thus cause death. In the museum of Owens College there is a specimen taken from the body of a man who died under the following circumstances. The man, when not quite sober, was turned out of a public-house, he tried to force his way in again and in doing so his finger was caught in the door, shortly afterwards he was found dead in the street. The body was taken to the Manchester Royal Infirmary, and Professor Young, who was then the pathologist, made a post-mortem examination, but found no cause of death until on opening the larynx a small piece of skin was discovered just below the glottis, further examination of the body showed that the fragment of skin had been detached from the index finger of the right hand. What probably happened was that on receiving the injury the man put his finger in his mouth to ease the pain, and the piece of skin in question, being all but detached, was carried away by a quick inspiration and lodged between the vocal cords.

**Homicidal suffocation** from the introduction of **foreign bodies** into the air-passages is rarely met with, except in infants or young children. In the case of children, various substances have been forced into the air-passages such as rolled-up pieces of cloth, pieces of newspaper, sand, earth, and artificial teats—the last-named to give colour to the assertion that death was accidental. Adults murdered by the introduction of foreign bodies into the air-passages have generally been under the influence of drink at the time or have been aged and feeble persons. Littlejohn<sup>1</sup> records the case of a woman who died suddenly, and at the post-mortem examination a wine-bottle cork (most probably thrust into the throat when the woman was intoxicated) was found covered with frothy mucus and tightly inserted into the upper part of the larynx, the sealed end of the cork with the mark caused by the corkscrew was uppermost. The defence was that the woman had drawn the cork from a bottle with her teeth and that it was propelled down the throat accidentally, the fact of the sealed end being uppermost, however, disproved this, and the frothy mucus which covered the cork indicated that it was introduced during life. The act was, doubtless, homicidal, but the accused was not found guilty, the Scotch verdict of “not proven” being given.

**Suicidal suffocation** from the introduction of **foreign bodies** into the air-passages is limited to lunatics. A woman in an asylum, aged thirty-eight, was found early one morning dead in bed with part of a stocking protruding from her mouth, death having resulted from suffocation, no noise nor disturbance nor anything wrong was noticed by other patients who slept in the same ward. At the inquest a long stocking was exhibited which had been removed with great difficulty from the air-passages.<sup>2</sup> Sankey<sup>3</sup> mentions the case of a male epileptic who committed suicide in an asylum, and was found dead lying on his back in bed with a round pebble in each nostril, and a strip of flannel rolled up and stuffed into the throat.

(b) **Suffocation from forcible compression of the chest** may be **accidentally** occasioned when large numbers of people are massed together on occasions of public rejoicing, or in panics, such as follow an alarm of fire raised in a crowded

<sup>1</sup> *Edin Med Journ*, 1885

<sup>2</sup> *Brit Med Journ*, 1882

<sup>3</sup> *Brit Med Journ*, 1883

building. A lamentable instance occurred in 1883 in Sunderland. Some hundreds of children in the gallery of a public hall rushed downstairs and were arrested by a closed door, a block took place at the bottom of the stairs and about 300 children were piled up so as to form an inextricable mass some 7 or 8 feet high. About 200 were killed, for the most part from compression of the chest, although in very few instances were the ribs fractured, which may be accounted for by the flexibility of juvenile bones, and by the absence of powerful struggling, such as takes place in a crowd of adults. Lambert,<sup>1</sup> who examined many of the bodies, states that the appearances after death were almost uniformly—congested puffy face, purple or blackish turgescence of the vessels of the neck, eyelids closed, eyeballs protruding, pupils dilated to the uttermost, and froth surrounding the mouth and nostrils, in twenty-four hours much of this had passed off, the face presenting the appearance of peaceful repose. Cadaveric rigidity was universally absent. In almost all the cases urine and fæces had been expelled.

**Suffocation of Infants in Bed.**—Infants are occasionally found dead from suffocation, while in bed with their parents or others, but it is not easy to determine in every case the exact circumstances of the death. What undoubtedly happens in some cases is that the infant is placed with its head on the pillow but slips down. The mother in her sleep pulls the bed-clothes around her neck and over the head of the child, thus suffocating it. In other cases the mother takes the infant in her arms for warmth or to suckle it, then goes to sleep, and the child is suffocated by pressure against the breast. This cause of death is spoken of as "overlying."

The post-mortem appearances are those of asphyxia. Signs of pressure such as flattening of the nose may be present, but in the Editor's experience, are exceptional.

In 1919, the deaths of 473 infants (258 males, 215 females) were recorded as due to suffocation in bed in England and Wales. Of this number, 195 were under one month and 466 under one year of age. The annual number of these deaths has declined steadily and rapidly during recent years, having been as high as 1,157 in 1911. A degree of uncertainty necessarily attaches to the circumstances and cause of death in infants found dead in bed with adults when the post-mortem appearances show only signs of suffocation, and there is reason to believe that a considerable proportion of deaths attributed to overlying are really due to natural disease. There are several pathological conditions, the post-mortem signs of which are practically indistinguishable from those of asphyxia, such as rickety convulsions and conditions somewhat vaguely certified as infantile debility. Even in the early stages of broncho-pneumonia, the patches of consolidation may be overlooked at the post-mortem examination unless this is made by someone well versed in pathological appearances. In a young infant, the parents may have failed to notice that it had been ailing, or may have attributed such symptoms as it showed to a trivial cold. The Editor has elsewhere examined this question in detail.<sup>2</sup> Briefly, the reasons for the above conclusions are that the relation between deaths from overlying and overcrowding is not constant, the urban mortality always being much higher than that in rural districts, though the overcrowding in the latter when expressed in terms of occupants per room is often nearly as high as that in urban districts, further, there is a marked seasonal variation in deaths from overlying, the mortality being highest in winter and lowest in summer. In both these respects

<sup>1</sup> *Brit. Med. Journ.*, 1883.

<sup>2</sup> *Statistics of Death from Violence and Unnatural Causes*, 1915.

the mortality from overlying behaves in the same way as the mortality from broncho-pneumonia, infantile convulsions, and infantile debility. The most conclusive argument, however, is to be drawn from the experience of one of the London coroners, who, being dissatisfied with post-mortems made by general practitioners, required all his post-mortems for several years to be made by expert pathologists. During this period his verdicts of overlying were less than 1 per cent of the total inquests on children under one year, whereas in the other coroners' districts the figures ranged from 8 to 25 per cent. In view of the difficulties of distinguishing the post-mortem appearances of overlying from those of certain forms of disease, the facts here mentioned should not be regarded as a reflection upon the general practitioner or as an argument that he should not (except in special cases) perform post-mortems for coroners' inquests.

It has been stated that drunkenness of the mother is largely responsible for deaths from overlying. In the Editor's experience after hearing a large number of these deaths enquired into, this factor has rarely been present. In support of the view that the ebriety is an important cause, it has been pointed out that the proportion of these deaths which occur on Saturday nights is larger than on other nights of the week. The Registrar-General specially investigated this question in his report for 1916, and showed that there is some preponderance on the Saturday night, and he appeared disposed to accept the view that this was due to drunkenness of the parents. It must be remembered, however, that among the working classes early rising is necessary during the week, and on Sunday they can prolong their rest. Thus, there is a longer interval during which the accident of death of the child in bed with the parents may occur.

The Editor is of opinion that the decline in infant mortality from overlying during recent years, to some extent at least, has been due to greater appreciation among practitioners of the possibility of natural death in these cases, and more minute investigations at post-mortems. In France deaths from overlying are almost unknown and in that country the autopsies in medico-legal cases are performed by expert pathologists.

**Homicidal suffocation from compression of the chest** has been effected in new-born infants. In adults, unless previously rendered incapable of resistance by intoxication, or by debility from age, such a mode of homicide would be scarcely practicable. The oft-quoted cases of Burke and of Bishop and Williams were examples of combined suffocation, they compressed the chests of their victims by allowing the weight of their bodies to rest on them, and at the same time covered the mouth and nostrils with the hands. Maschka met with the case of a girl of fourteen who was suffocated by pressure on the chest by one man, whilst another perpetrated a rape upon her.

(c) **Suffocation by covering or compressing the mouth and nostrils** is a mode of suffocation usually designated **smothering**. In the case of infants it may easily occur either **accidentally** or with homicidal intent by simply covering them over with a thick layer of clothing, such as the bed-clothes, mere continued pressure of the child's face against the mother's breast in the act of suckling has proved sufficient. Adults may be accidentally smothered by immersion in substances composed of loose particles, such as grain, sand, flour, and the like. Falk<sup>1</sup> records the case of a man who was accidentally buried in a large quantity of flour. The mouth and œsophagus were filled with flour agglomerated with saliva, but none obtained access to the stomach, the lungs and the

<sup>1</sup> *Nordiskt med. Arkiv*, 1891.

brain were hyperæmic, particles of flour were found in the bronchi as far as the smallest ramifications. Russo-Gilberti and Alessi<sup>1</sup> found loose particles of the surrounding material in the larynx, bronchi, and even in the pulmonary alveoli of animals that had been buried alive. Epileptics are not unfrequently smothered during a seizure, either when in bed, from close approximation of the bed-clothes to the face, or in the daytime from falling on to some yielding substance face downwards. Janeway records the case of an epileptic who had an attack in a stable-yard and was suffocated by the manure on which he fell, portions of which were found in the larynx after death.

**Homicidal suffocation by smothering** is, doubtless, of frequent occurrence as regards infants and young children, the modes which have been adopted are described in the section on infanticide.

Adults have either to be taken unawares or to be asleep or intoxicated to permit of death being homicidally produced by smothering. Plasters made of pitch or other sticky substances have been used for the purpose of rendering persons helpless and incapable of raising an alarm whilst being robbed, if the plaster is not quickly removed death from suffocation results. Wald<sup>2</sup> records an instance of homicidal smothering of an adult. A man resolved to murder his wife, and took the opportunity whilst she was asleep in bed to cover her over with several heavy bed-covers and then laid himself on the top, in a short time, the struggles she at first made ceased, and the woman was dead. The man then informed the doctor that his wife was attacked with cramps and desired his attendance, but before the doctor arrived he met him with the intelligence that she was dead, no suspicion was created, and the body was buried. Information was subsequently given to the authorities as to the real cause of death, and the body was exhumed and examined. Signs of death from asphyxia were present, and the murderer subsequently confessed to having perpetrated the crime in the way above described.

**Suicidal smothering** is almost unknown. A case<sup>3</sup> occurred in France in which a woman got under the bed-clothes after desiring her little child to bring all the cushions, clothes, and other similar articles that were in the room and pile them on top of her, the child did so, and some hours after when the woman was discovered she was dead. Signs of death from asphyxia were found at the necropsy.

The modes in which suffocation can be produced being so varied, the **post-mortem signs** within certain limits are correspondingly diversified.

**Externally**, the appearances comprise all degrees of significance from those cases in which the mouth and nostrils are found filled with a foreign substance, down to those in which death has resulted from smothering by graduated pressure with a soft material, the first being obvious at a glance, the second yielding no external trace whatever of the cause of death. Apart from any special indication afforded by the presence of injuries about the mouth or nostrils, or of some of the material by means of which the air-passages were obstructed, one or more of the general signs of death from asphyxia may be visible, such as punctiform ecchymoses on the conjunctivæ and on the outer surface of the lower eyelids. Such ecchymoses are not to be accepted as proof of death from suffocation, they are occasionally produced by epileptic attacks, Cabade<sup>4</sup> relates the case of a healthy man, aged forty, who, in consequence of an epileptic seizure, developed numerous subcutaneous extravasations of blood on the thorax and abdomen. They may even be produced after death.

<sup>1</sup> *Archivio per la Scienze Mediche*, 1888

<sup>2</sup> *Gerichtliche Medicin*, 1858

<sup>3</sup> Wald, *l.c.*

<sup>4</sup> *The Lancet*, 1892

Haberda,<sup>1</sup> by suspending, head downwards, the bodies of infants shortly after death developed ecchymoses under the conjunctivæ indistinguishable from those due to suffocation. Turgescence of the face and large vessels, with a cyanotic hue of the extremities is suggestive of suffocation from compression of the thorax, but the condition disappears in a few hours after death.

**Internally**, there may or may not be the usual signs of death from asphyxia, the amount of blood contained by the lungs and heart is by no means constant, nor is the dark colour and fluidity of the blood invariable. In children, especially, an œdematous condition of the lungs is often found after death from suffocation. Sub-pleural ecchymoses, which Tardieu erroneously regarded as exclusively due to asphyxia produced by suffocation, are usually present, the pericardium also may be ecchymosed. The degree of hyperæmia of the other organs—brain, stomach, intestines, spleen, liver, and kidneys—is subject to great irregularity. It is to be remembered that the **absence** of the so-called **signs of death** from **asphyxia** does not exclude the possibility of death having resulted from suffocation.

In the case of an **infant** alleged to have "**died in a fit**"—the circumstances being suspicious of accidental overlaying or of criminal suffocation—too much importance must not be attached to the presence of sub-pleural ecchymoses. The ecchymoses are, to a great extent due to increased arterial blood-pressure, therefore a convulsion is likely to produce them, and further, since both in convulsions and in suffocation death is mostly due to asphyxia, the **post-mortem** appearances respectively found in such cases may be very similar. (See p. 163.) Occasionally, however, after death from convulsions, the post-mortem appearances of death from asphyxia may be absent. Sub-pleural ecchymoses have even been found under circumstances which placed suffocation out of the question, as in the instance mentioned by Degranges<sup>2</sup> in which they were present in an infant at term, that was removed by Cæsarean section, without being subjected to any possibility of suffocation.

When making an inspection of the dead body for medico-legal purposes, the **air-passages** from the mouth down to the bronchi should **always** be examined, otherwise the real cause of death may remain undiscovered. It is too frequently assumed, if the conditions of the heart, the lungs, and the blood usually associated with death from asphyxia are absent, that there is no need to examine minutely into the state of the respiratory tract, more than once it has occurred that a medico-legal inspection which yielded negative results as to the cause of death has been proved to have been negligently made, by the discovery of a foreign substance in the air-passages on subsequent and more thorough examination. In addition to the detection of foreign bodies in the air-passages the state of the mucous membrane of the larynx and trachea requires investigation, it is sometimes found injected, and may be covered with frothy mucus, possibly blood-stained, this would point to an asphyxial mode of death, though its absence would not justify the opposite conclusion.

All signs of general violence, such as bruises, scratches, or ruffling of the epiderm, are to be carefully looked for. If the examination is made before the body is moved, the surroundings may afford valuable indications, therefore careful inspection of every detail must be made and *notes taken before it is disturbed*. It not unfrequently happens that conditions subsequently discovered may receive important corroboration from some apparently trifling detail which, without this precaution, would have escaped notice, or from want of aid to the memory afforded by the notes would probably be imperfectly interpreted.

<sup>1</sup> *Internat Med Congress*, 1897.

<sup>2</sup> *Gaz des Hôpitaux*, 1867.

### DROWNING.

**Drowning** is a mode of death from asphyxia caused by continuous or by intermittent submersion of the mouth and nostrils under water or other fluid, so that access of air to the lungs is either at once or gradually cut off until life is extinct. When asphyxia is caused by hanging or by strangulation, access of air to the lungs is simply obstructed, when it is caused by drowning, an irrespirable medium is drawn into the air-passages by attempts at respiration, and in this way special changes are produced in the lungs beyond those which are due to uncomplicated asphyxia, these changes constitute one of the principal signs of death from drowning. Death may result from falling into the water and still not from asphyxia—fatal syncope may occur at the moment of submersion, or in rare instances the head or abdomen may strike against a rock or other solid body with sufficient violence as to cause immediate death from shock—in such cases the characteristic signs of death from drowning will be absent.

From experimental investigations Brouardel and Loye<sup>1</sup> consider that the mode of death from submersion usually attributed to syncope is due to inhibition, and they emphasise the distinction between the manner in which the respiratory act is influenced by drowning on the one hand and by external constriction of the air passages on the other. When asphyxia is produced by hanging, the entire muscular force of the respiratory apparatus is thrown into immediate action in the endeavour to obtain oxygen. When submersion is the cause of asphyxia, there is at first an attempt on the part of the animal to breathe as usual, then appreciating the danger, it struggles to escape and at the same time fixes the thorax voluntarily suspending respiration in order to prevent the water from entering the air passages, the respiratory centres, however, are soon stimulated beyond the control of volition, and involuntary respiration movements are set up by which water is drawn into the lungs and penetrates the alveoli. In addition to the resistance offered by the will, the sensory nerves of the skin and of the air passages play an important rôle.—If the trachea of an animal is opened and a tube is inserted in the opening, the glottis has no longer control of the air way, and it might be supposed on submerging the animal that water would at once be drawn into the lungs, the same suspension of respiration, however, takes place as before, from fixation of the thorax, so long as the centres can be kept in abeyance. But if both pneumogastrics are divided high up in the neck, and a tube placed in communication with a vessel of water is introduced into the trachea, the animal continues to make the usual movements of respiration for thirty seconds, although water is being received into the lungs in place of air, the animal then becomes agitated, but still continues the respiratory movements. If, after division of the pneumogastrics, an animal is plunged in water it fixes the thorax almost as though the nerves were intact, thus demonstrating that the sensory nerves of the skin and of the naso-pharyngeal tract take part in the inhibition of respiratory movements, in the preceding experiment they were not called into play. When an animal chloroformed until the cornea is insensative is submerged, it makes no attempt to prevent water entering the air passages, but continues the movements of respiration as before submersion. From these experiments Brouardel and Loye infer that, although volition has to do with the temporary arrest of respiration, irritation of the nasal and laryngeal nerves exercises a still more powerful influence by inhibiting respiratory movements. A transient stage of interrupted respiration may exceptionally be replaced by one of permanent inhibition, and death from submersion is thus caused with absence of the usual signs produced by the presence of water in the lungs.

When death takes place from drowning in the ordinary way, submersion is usually continuous—even when the individual falls from a height above the water-level, and in consequence of the acquired momentum reaches a considerable depth, if not prevented in some way he generally returns at least momentarily to the surface. This is not due to the body being of less specific gravity than water, but to involuntary movements of the limbs—apart from skilled movements, such as are executed by practised swimmers. The specific

<sup>1</sup> *Archives de Physiol. norm. et pathol.*, 1889.

gravity of the human body is slightly greater than that of water, being about as 1.8 is to 1.005 or 1.03, according to the kind of water—fresh or salt. All the constituents of the body are specifically heavier than water, except fat, and lungs which have breathed, and these do not suffice to counterpoise the heavier tissues, still, the difference is so slight that very little movement brings the body to the surface, if only for a moment. The greater the percentage of fat and of lung-capacity in relation to the body-weight, the greater the tendency to float, women, as a rule, float better than men because of the slighter build of the skeleton and large proportion of fat (see p. 46), new-born infants, if well nourished, will scarcely sink unless weighted.

The body of a person who has died from drowning is deprived of the buoyancy of the lungs, since the air they contained has been almost entirely replaced by water, this, together with the absence of movement, causes the body to sink and to remain submerged. Submersion continues until the gases of putrefaction are developed in sufficient quantity to reduce the specific gravity of the body below that of the surrounding medium, the body then rises to the surface unless prevented by some mechanical obstruction—such as being covered over with sand or mud, or being caught by weeds, ropes, or the like. In temperate climes flotation generally occurs within the first week, the exact period after death depends upon the initial difference in specific gravity between the body and the medium in which it is submerged—chiefly determined by the amount of fat—by certain intrinsic conditions depending on the degree of vital exhaustion that preceded death, and on the temperature and stillness of the water. A body submerged in deep water will not rise so soon as one lying in shallow water, because the latter will be warmer and will consequently promote an earlier formation of the gases of putrefaction. A body submerged in a pond is likely to rise earlier than one at the bottom of the river, because the still water acquires more heat from the sun's rays and from the stratum of air above it than the flowing water of a river.

**Post-mortem Appearances.**—When the body of a person who has been drowned is removed from the water within a few hours after death and is examined forthwith, there is usually little difficulty in determining that death was caused by drowning, on the other hand, when the body remains in the water until decomposition is advanced, insuperable difficulties are interposed which may render the formation of anything like a positive opinion impossible. The interval that elapses between the removal of the body from the water and the examination largely influences signs of great diagnostic value.—The froth visible about the mouth and nostrils in the recently drowned is soon dissipated on exposure to the air, the surface of the body—especially if the clothing is removed—in a short time becomes dry and discoloured, putrefactive changes in the body advance with great rapidity after its withdrawal from the water, and in a comparatively short time completely obliterate the indications which distinguish death by drowning from other modes of death with subsequent submersion of the body. The following description refers to the bodies of those recently drowned and examined shortly after removal from the water—

**External Appearances.**—Rigor mortis usually comes on early, in some cases immediately, so that the last position of the limbs during life is maintained after death. It is not always easy to determine whether or not rigor mortis is actually present, as in cold water a stiffening due to solidification of the body fat occurs which may persist in the water long after true rigor mortis has passed off. This condition rapidly disappears when the body is placed in a warmer environment. The surface is pallid, but not more so than of a dead body.

under ordinary conditions. The face is tranquil and the eyes and mouth are partially open. There may be rosy patches on the face and neck, and in some instances the face is of a reddish-blue or violet hue, at a later period after death, its colour may be changed to a dirty red. The skin often presents the appearance known as "goose-skin," a condition met with after a variety of sudden deaths by violence, it is due to vital reaction, the manifestation of which persists after death from the occurrence of instantaneous cadaveric rigidity in the *arrectores pilorum* when in a state of contraction. The skin of the palms of the hands, the soles of the feet and the knees is bleached, sodden, and wrinkled, the result of imbibition, it is of no significance other than showing that the body has lain in the water for twelve or more hours. The tongue has been described as being pressed forward between the teeth, it is only exceptionally that this occurs, and as the same condition is found after violent death from other causes, it is a sign of little import. The **most valuable** of the external indications that death has resulted from drowning is the presence of a **fine froth** on the lips and nostrils. This froth or foam may be white or it may be blood-stained, it has been not inaptly likened to the finely constituted lather produced by shaving-soap. It is probably found after drowning under all circumstances and in all media, and has been in the case of an infant drowned in a cesspool. Its continuance, however, cannot be depended on for more than four days in winter and about three in summer. The body being under water, if it has disappeared from the lips, a little pressure on the chest may cause some to well out of the mouth. The presence of substances, such as weeds from the bed of a river, or of fragments of clothing, clutched in the fingers—due, in the first instance, to a vital act subsequently rendered permanent by instantaneous cadaveric rigidity—is evidence of submersion during life, in practice it is a sign not frequently met with.

**Internal Appearances.**—The **lungs** usually exceed their normal volume, so that they almost cover the pericardium, a condition sometimes described as "ballooning" of the lungs, instead of collapsing when the thorax is opened, they protrude so as to fill the aperture made by the removal of the sternum. Care must be taken not to confound this condition with distension of the lungs due to gases of putrefaction. The consistence of the lungs is doughy—they retain the pit produced by pressure of the finger. They are sometimes of a pale grey, with reddish stains produced by transudation of water tinged with blood-colouring matter, in other instances the entire surface of the lungs is of a reddish-blue. Draper, in the majority of cases examined by him, observed punctiform subpleural ecchymoses, though few in number and mostly at the lower part of the lobes. Most observers state that they are of very exceptional occurrence in death from drowning, Ogston met with them in barely 7 per cent of the cases he examined. On section, the lung-substance is found to contain water which, along with froth, exudes from the cut surface on the slightest pressure, the bronchi, and possibly the alveoli, contain fine froth of the same character as that on the lips and nostrils, the red stain, if visible on the surface of the lungs, is seen to pervade their entire substance.

Two views are held with regard to the condition of the parenchyma of the lungs after death from drowning. According to one, it is infiltrated with water or other medium in which drowning takes place, and this to a certain degree accounts for the increased volume of the lungs. According to the other view, there is but little fluid in the parenchyma, and any that is present is derived from exudation from the vessels producing oedema. Paltauf<sup>1</sup> states that the water which is drawn into the lungs in attempts at respiration enters the

<sup>1</sup> *Ueber den Tod durch Ertrinken*, 1883



alveoli, and finds its way from thence by the lymph spaces and occasionally through small lesions in the alveolar walls, into the interstitial alveolar, subpleural, peribronchial, and perivascular connective tissue. Lesser,<sup>1</sup> on the other hand, states that a secretion of mucus, caused by aspiration of water into the air passages, blocks the finer bronchi so that water cannot reach the alveoli except in scarcely appreciable quantity, he denies that the parenchyma contains water, if any fluid is there it is derived from the blood vessels and is a vital exudation product—a true œdema. It is probable that a mucous secretion in the bronchi does occur during death from drowning, but only in quantity when the death struggle is prolonged. Experiments on animals show this, and also that blocking of the finer bronchi by the mucus is one reason why the lungs after death from drowning do not collapse when the thorax is opened—the alveolar contents cannot escape.

The fine froth contained in the air-passages is composed of a variable mixture of the drowning medium with air, mucus, and possibly blood. When the death-struggle has been prolonged more froth is produced, and it is of a more lasting nature than when the death occurs quickly, this results from the presence of a greater proportion of mucus, which imparts increased tenacity to the bubbles, and from more vigorous admixture of air, water, and mucus, which augments the total amount of froth. According to the observations of Brouardel and Vibert,<sup>2</sup> staining of the froth is due to infiltration of blood-colouring matter derived from small extravasations in the parenchyma of the lungs.

If, after submersion, the body does not return to the surface until death has taken place, there will be less froth and more water in the air-passages than when the mouth and nostrils momentarily emerge once or more and additional air is inspired, this is supposing that death takes place from asphyxia, if from syncope or, what comes to the same thing, from permanent inhibition of the respiratory movements, no froth at all will be present. The pleural cavities usually contain a quantity of water with or without a certain admixture of mucus. The **mucous membrane** of the **trachea** may be injected, but this sign is more frequently absent after drowning than after other modes of death from asphyxia, the lumen of the trachea is usually filled with froth, which may reach the mouth unless the body has lain for some time in water or is not examined soon after withdrawal from it. The **heart** usually presents the appearance due to death from asphyxia, the left side is empty or nearly so, and the right side filled with blood. The vascular condition of the **brain** and its membranes is not constant, and therefore is of no diagnostic value as regards death from drowning, Draper, in 149 examinations, found that injection of the membranes with hyperæmia of the brain was present in about half the number of cases. The **blood**, dark in colour and containing but little oxygen, is diluted with water, sometimes to the extent of one-third or one-fourth of its total weight (Brouardel and Vibert). The dilution is greater in the left side of the heart than in the right side, because the blood arriving by the pulmonary veins has received a fresh addition of water in passing through the lungs, the blood in the portal vein may also be much diluted from imbibition by its branches coming from the stomach, which, as will presently be explained, usually contains a quantity of water. The dilution of the blood is proportional to the duration of life after submersion—slow drowning produces much dilution, quick drowning less, when death from submersion results from syncope, or inhibition of respiration, the blood does not contain any excess of water, unless the body remains long submerged. After death from drowning the blood is usually *fluid*, as is the case in other modes of death from asphyxia. Brouardel and Loye found that when the body is examined soon after death from rapid

<sup>1</sup> *Atlas d'ger Med*, 1891, and *Vierteljahrsschr f ger Med*, 1884

<sup>2</sup> *Annales d'Hygiène*, 1880

drowning the blood for the most part is coagulated, but if an interval of one or two days elapses before the examination is made, the blood is again fluid Coutagne<sup>1</sup> and others deny that the blood coagulates, stating that it is fluid even when the body is examined shortly after death, a few dark clots may be found, but in feeble proportion to the mass of blood

There is nothing distinctive in the condition of the mucous membrane of the **stomach**, its vascularity being variable, occasionally ecchymoses may be seen, as in other modes of asphyxial death, and fine froth may be found either in the stomach or œsophagus In another respect, however, the stomach and the intestines yield valuable evidence as regards the occurrence of death from drowning When submersion in the living takes place, a portion of the water drawn into the mouth is frequently involuntarily swallowed, hence the presence of water in the stomach has been regarded as a sign of death from drowning In estimating the value of this sign, two questions naturally present themselves (a) Is water invariably found in the stomach after death from drowning? (b) Is it possible for water to find its way into the stomach of a submerged dead body? Answers to these questions have been sought for by experimental investigations on animals, and by statistical observations of human bodies in which death took place from drowning

(a) Experiments on animals drowned in water artificially coloured show, for the most part, that water is swallowed in the act of drowning In sixteen experiments by Fagerlund<sup>2</sup> some of the coloured water was invariably found in the stomach after death Misuraca<sup>3</sup> almost always found water in the stomach of animals that had been drowned

Observations made on human bodies as to the presence of water in the stomach after death from drowning yield less decisive results Some observers state that water in characteristic amount is almost invariably present, others found it in a limited number of cases, and often in such small quantities as to be worthless from the diagnostic standpoint Amongst recent observers, Hofmann<sup>4</sup> found the amount to vary considerably, only exceptionally was water present in bulk Ogston<sup>5</sup> found it present in 74.6 per cent of cases Tourdes<sup>6</sup> out of 93 cases found a considerable amount of water in the stomach in 37, but little in 34, and none at all in 22 Draper in many of 149 cases states that the stomach was either empty or contained remains of food with a very small quantity of fluid Lesser out of 30 cases found no appreciable amount in 9, a variable amount in 14, a layer of water distinct from the other contents of the stomach in 3, and in 4 water alone or mixed with a little mucus Paltanuf frequently found water in the stomach after death from drowning

(b) Obolonsky<sup>7</sup> placed in coloured water the bodies of 18 children of from two weeks to two months old, weighted so that they could not float, they were allowed to remain submerged for from one to three days In three of the bodies a considerable quantity of the coloured water was found in the stomach, in two others only a small quantity—these five bodies had lain seventy-two hours in the water, in the remaining thirteen no trace of water was found in the stomach Misuraca never found water in the stomachs of animals submerged after death Fagerlund experimented with a large number of dead bodies of children and animals by placing them in various positions—on the back, on the side, face downwards, etc.—under water, in some cases the mouth was kept open with pieces of wood, and the tongue was drawn forward and fixed, the results showed that water finds its way with great difficulty into the stomach after death Bougier<sup>8</sup> placed the dead bodies of twenty-three human beings and seventeen animals in various coloured fluids without water entering the stomach in any one case

From these experiments and observations the answers to the two questions will be—(a) That water is not invariably found in the stomach after death from drowning (b) That it is possible for water to find its way into the stomach of a dead body lying in water

<sup>1</sup> *Archives de Physiol norm et pathol*, 1891

<sup>2</sup> *Vierteljahrsschr f ger Med*, 1890

<sup>3</sup> *L'assistenza meccanica e le sue varie forme*, 1888

<sup>4</sup> *Lehrbuch der ger Med*, 1887

<sup>5</sup> *Edin Med Journ*, 1882

<sup>6</sup> *Dictionnaire Enc des Sciences Med*

<sup>7</sup> *Vierteljahrsschr f ger Med*, 1888

<sup>8</sup> *Thèse*, Paris, 1885

The amount of water found in the stomach is to be noted, when small it is of doubtful import, in some cases clear water is found in a separate layer above the other contents of the stomach. The occurrence of fluid in the stomach possessing special characteristics corresponding with the medium in which the body is found may be of importance, but it is not a proof of submersion during life. When submersion is followed by death from syncope no water is found in the stomach. **Absence of water** in the **stomach** after death does *not* exclude drowning as the cause of death.

In some cases a portion of the **water** that has entered the stomach during death from drowning is forwarded by vital contraction into the **intestines**, and its presence there has been regarded as almost certain proof of submersion whilst living. The questions put in relation to water found in the stomach are equally relevant in the case of the intestines. Fagerlund's very comprehensive experiments as to the conditions under which water may reach the intestines afford valuable evidence. The outcome of them is that, occasionally, during death from drowning, peristaltic movements cause water to pass from the stomach into the bowels, that the pylorus offers a certain amount of resistance to the onward passage of the water, that the water passes into the bowels more easily from an empty stomach than from one filled with food, and that slow drowning appears to favour its ingress. In the dead body water only finds its way from the stomach into the bowels when excessive pressure is brought to bear.

As premised, the signs of death from drowning described in the preceding pages are those found in recently dead bodies which are examined shortly—within twelve or, at the most, twenty-four hours after removal from the water, in warm weather even this interval will be sufficient to obscure the more significant indications. When putrefactive changes are advanced beyond the early stages, the frothy contents of the bronchi and trachea will have disappeared—there may be fluid and a number of air-bubbles, but no lathery froth. At a certain stage of putrefaction the lungs collapse on opening the thorax, or are already collapsed, in the former case, the softening undergone by the lung-tissue and by the mucus of the finer bronchi permits the alveolar contents to escape under the pressure of the atmosphere, in the latter, the pressure will have been already produced by transudation of water into the pleural cavities. The colour-changes, both external and internal, resulting from putrefaction are very deceptive, especially in relation to the possible occurrence of bruises, contusions, and the like, so that after the withdrawal of a body from water in which it has lain for twenty-four hours or more, bruises, though present, may not be visible on account of imbibition by the skin, when the skin dries they become apparent.

The longer the body has been submerged the less valuable is the presence of water in the stomach or intestines as a sign of death from drowning. Water, mud, and particles of sea-weed may find their way to a limited extent into the air-passages of a body submerged after death, grains of sand have been found even in the alveoli, but there is no great increase in the volume of the lungs, nor is fine froth present in the bronchi. Transudation of water into the pleural cavities may buoy up the lungs in the dead body and make them look prominent on removal of the sternum without there being any "ballooning."

Exceptional cases occur in which, after a very short period of immersion in water containing substances of an irritant nature, death results from asphyxia due to a subsequent rapid exudation of mucus into the air passages. Brockbank<sup>1</sup> reports a case illustrative

<sup>1</sup> *Med Chron*, 1894

of this mode of death. A man, aged thirty nine, accidentally fell into a canal at a spot where barges carrying carboys of acid are unladen, a carboy not infrequently being broken and its contents spilt into the canal. The man was rescued in less than a minute and appeared to be little the worse, but in about half an hour he became "stuffed," and soon after was seized with "fits," in one of which he died in less than an hour after immersion. At the necropsy, white froth, along with blood stained fluid, was found about the mouth, both lungs were cedematous and the bronchi were filled with blood stained frothy serum, the mucous membrane of the larynx, trachea, and bronchi being congested. The mode of death and the post mortem appearances resembled those produced by inhalation of acid fumes, with the exception that the period of survival was shorter.

**Epitome of the Important Signs of Death from Drowning.**—The only external sign of import is the presence of a **fine froth** or lather on the lips and nostrils, this sign is of great significance, but its duration after death is limited. An adventitious sign constituted by the presence of weeds or other local objects grasped in the hands is also indicative of submersion during life, but its occurrence is not frequent.

The internal signs are **increased volume** of the **lungs** with doughy consistency, water in the pleural cavities, exudation of water and froth on section of the lungs, **fine froth** in the bronchi and trachea, dilution of the blood, and **water** (or the fluid in which the individual was drowned) in the **stomach**, especially if some has reached the **intestines**.

In the event of an investigation made on a dead body removed from water indicating death from drowning, the next question is Was drowning the result of **accident, homicide, or suicide**?

First, as to probabilities. For the year 1919 the deaths from drowning in England and Wales are thus apportioned in the Registrar-General's reports

Accident		Homicide		Suicide	
Males	Females	Males	Females	Males	Females
1,537	341	16	11	425	349

These figures show that by far the largest number of deaths from drowning are due to accident, and male sex contributing more than four times as many as the female sex. This is owing to occupation risks, which affect men much more than women. Of the 27 deaths from homicidal drowning, in the periods examined, 22 occurred in children under two years of age. Under ordinary circumstances it is difficult to murder an adult by drowning, unless the victim is in some way rendered incapable of resistance, with women the difficulty is not so great as with men, because women are more helpless when unexpectedly plunged into water.

When suicidal is compared with accidental drowning, the sexes show a marked numerical alteration, the proportion of females to males being about as 5 is to 6. With women, drowning is the preferred mode of committing suicide, probably because it requires no mechanical contrivance for its accomplishment, the simple act of falling into water is sufficient.

As a means of ascertaining whether drowning was due to accident, homicide, or suicide, the external relations are of increased importance as compared with those attending other modes of death from violence. If the body is clothed in the usual manner, the condition is consistent with any one of the three,

but if the body is without clothing, and the signs of death from drowning are present, either accident or suicide is indicated- the former if the place and the season of the year are suitable for bathing, the latter, when one or both are unsuitable. A hasty conclusion is not to be arrived at, even when circumstances seem exclusively in its favour, and, above all, no opinion should be expressed until a full examination of the body has been made. Illustrative of the necessity of this is a case related by Winsor<sup>1</sup>. The body of a woman was found naked in water about two feet deep, at first it was thought to be a case of accidental drowning, as the woman had been heard to express a desire to bathe in this particular place. On examination, however, an extravasation of blood was found in the right pectoral muscle, but none of the usual signs of death from drowning were present- no water was found in either lungs or stomach. The conclusion arrived at was that death had been caused by suffocation from pressure on the chest (probably by some one kneeling on it), and that the dead body had been placed where it was found.

**Bruises and wounds** on the body give rise to suspicion of foul play, but they may result from previous accident or suicidal attempt, or from injuries produced after death by the body being tossed against rocks, posts, or other objects by the current or movement of the water. Draper<sup>2</sup> relates two instructive cases of antecedent accidental injuries which complicated the diagnosis as to the mode of death. The dead body of a brewer was found in a large reservoir of water in the basement of a brewery, the sole access to the reservoir was through a shaft, barely two feet square in transverse section, the opening of which was in an upper storey of the building. The body showed undoubted signs of death from drowning, along with contusions on the face and top of the head, which did not seem likely to have been produced by falling down the shaft- it looked as though the deceased had been rendered unconscious by a blow on the head and then thrown down the shaft. On inquiry it turned out that the bruises had been produced in a drunken brawl two days before death, and a letter was found on the body in which the deceased signified his intention to commit suicide. The second case was that of a man whose body was found in a river, with a lacerated and contused wound behind one ear which laid bare the skull and which had the appearance of having been caused, during life, by a sharp stone or other similar object, the signs of death from drowning were present. Here also the indications pointed to a blow on the head having been inflicted by some one, who afterwards threw the unconscious victim into the river. The body was ultimately recognised as that of a man who had fallen down a flight of steps while intoxicated, and who, being without means of sustenance, had subsequently committed suicide.

Injuries may be sustained in the act of jumping into the water, in this way fractures of the vertebræ, of the bones of the limbs, and also of the skull have been occasioned. Ogston states that in a woman who leaped over the parapet of a bridge, the perineum was lacerated by forcible separation of the thighs on coming in contact with the water.

As previously stated in relation to another mode of violent death, suicides, in order to terminate their existence, not unfrequently make more than one attempt, which may vary in kind- a man may put a bullet into his chest and then drown himself, or he may cut his throat before doing so. It is by no means an exceptional occurrence for a man or woman to take poison and then to throw themselves into water, such cases demand great perspicacity in their interpretation, for the unlikely frequently happens. For example, the combination

<sup>1</sup> *Boston Med and Surg Journ*, 1889

<sup>2</sup> *Boston Med and Surg Reporter*, 1885

of poison with drowning is suggestive of suicide, but it has been due to homicide. A woman took advantage of another woman's liking for brandy to give her some which she had previously mixed with arsenic, the brandy was drunk in the open air, and, whilst the victim was sitting on the bank of a river with her back towards the water her companion, losing patience at the slow action of the poison, pushed her in, the body being found three days after. Such a case is clearly beyond the scope of medical evidence, so far as determining whether death was due to homicide or suicide is concerned, the lesson to be learnt from it is, not to express a decided opinion on a probability, which in this instance was very great, since the combination of poison and drowning is so rarely the result of a homicidal act that Bělohradsky,<sup>1</sup> who relates the above case, only met with it twice in more than a thousand cases. Whilst the probability of such combination is in favour of suicide, the possibility of homicide must not be ignored.

Strangulation is an exceptional method of committing suicide, therefore, if a dead body is found in water with signs of strangulation and with absence of the signs of drowning, the probability is in favour of homicide by strangulation and subsequent disposal of the body in water. A case recorded by Hofmann<sup>2</sup> constitutes an exception to this statement. A girl, about twenty years of age, was found dead in a bath with a piece of thick pack-thread tied round the neck, the naked body was in a kneeling posture, the head being under water, examination showed that death resulted from strangulation. The bath was in a public institution, and the deceased locked herself in the bath-room, the door of which was under the observation of the attendant, there could, therefore, be no doubt that the girl had taken her own life. It might be inferred, had the case been one of homicide, that the murderer would have removed the evidence of strangulation before placing the body in the water, but Bělohradsky relates an instance which affords a further illustration of the risk of error which is inseparable from inferences founded on probabilities. A dead body, the neck of which was surrounded three times with a cord, was found in water, but no signs of death by drowning were present, the case was one of homicide by strangulation, the body having been placed in the water after death. Care must be exercised not to mistake marks produced on the neck at the time of or soon after death for indications of foul play, in one such case the string of a cloak, and in another the branch of a tree produced deceptive appearances.

Indications of severe blows on the head, having the appearance of injuries sustained during life, may give rise to suspicion of homicide, the suspicion is strengthened if the signs of death from drowning are absent. The possibility of the injuries being independently produced within a limited period before death has to be taken into account. Except in the case of young children, cut-throat wounds on bodies found in water are strongly indicative of suicide.

When a dead body is found in water with a wound caused by a fire-arm, attention must be paid to the position of the wound, the inference usually is, that death resulted from suicide, but, to make this probable, the wound should be on the front of the body, in the head or chest. If the body is found in still water— as in a pond—the firearm should be near it, if in a river or in the sea, the body may have floated to a considerable distance from the spot where the shot was fired. There is obviously nothing to prevent a murderer from leaving the weapon near the body in order to suggest suicide, but such

<sup>1</sup> Maschka's *Handbuch*

<sup>2</sup> *Wiener med. Presse*, 1879

a proceeding would be risky, unless he had possessed himself of a firearm belonging to the deceased

**Injuries produced after death** occur mostly in moving water—in rivers with swift currents, in harbours, or in the locks of canals. Sometimes the injuries strangely simulate in position and extent wounds such as are frequently inflicted during life, in other instances, the body is smashed in a way that is incompatible with even extreme homicidal violence. As a rule there is no difficulty, but care is occasionally necessary to distinguish between wounds inflicted at the time of death and injuries produced afterwards which co-exist on the same body. The characteristic appearance of wounds made before and after death are described in a succeeding chapter. When injuries to vital organs are found, which from their nature would necessarily be immediately fatal, and the body shows distinct signs of death from drowning there can be no doubt that the injuries were produced after death. On the other hand, the absence of the signs of death from drowning and the presence of injuries of this type would be suspicious of foul play, unless, from the nature of the surroundings, it might be supposed that the body had fallen into the water unaided, after the victim himself had inflicted the injuries—a man has been known to blow his brains out on the bank of a river and to fall dead into the water. A suicide who throws himself into the water from a great height, by striking some object may fracture and dislocate the spine in the cervical region, in such a case the signs of death from drowning would be absent, the manner in which death was brought about would have to be inferred from the surroundings. The possibility of death occurring from syncope on submersion with the subsequent production of injuries by the body being thrown violently against projecting objects must be borne in mind, as in this case there would be the combination of injuries, which, from their nature, might be expected to be immediately fatal, with absence of signs of death from drowning.

A case reported by Richardson<sup>1</sup> illustrates some of the points mentioned. The dead body of a man, of about 60 to 70 years of age, was found in the sea, decomposition was taking place, and was more advanced in the head than in the abdomen. The nasal bones were broken and loose, the scalp was torn away on each side of the head at the junction of the temporal and parietal bones, the bare skull was rough as though it had been rubbed against some hard substance, but the bone was not fractured. The spine was dislocated between the third and fourth cervical vertebrae, the spinous process of the latter being broken, and the cord crushed. The lungs were distended and spongy, they overlapped the heart, and when cut into exuded bloody froth. The blood was fluid and dark. The stomach contained a pint and a half of fluid, chiefly sea water. It was correctly inferred that death took place from drowning, and that the injuries were caused after death. The cardinal signs of drowning were found associated with an injury to the cervical cord which, had it been produced during life, would have been immediately fatal. Suicide was indicated by the month—November—together with the fact that the body had stockings on and also the wristband of a flannel shirt.

Occasionally a body presenting the signs of death from drowning is found in water only a few inches deep, such cases are usually suicidal, but they may be accidental, the deceased being intoxicated or under the influence of an epileptic seizure which caused him to fall into the water face downwards in a helpless condition. Except in the case of very young or very old subjects, it would be difficult to murder any one in this way, unless the victim was

<sup>1</sup> *Brit Med Journ*, 1889

previously rendered incapable. Bruises or superficial marks of violence on the body having the characteristics of those produced during life are to be interpreted with caution, as a drunken man may have previously fallen more than once and thus occasioned the injuries, or they may have been produced by clonic epileptic spasms.

Drowning has been caused by placing the head only under water contained in a tub or other receptacle, in the case of adults the act is usually suicidal, the exceptions are when helplessness, occasioned by previous intoxication or other causes, has been taken advantage of to commit homicide. Infants have been drowned in this way and their bodies afterwards thrown into a pond or river, in one such case, the fluid found in the air-passages consisted of dirty water containing soap, which demonstrated that the child was not drowned in the medium in which the body was found. Occasionally death may be due to accident as when a workman falls head downwards into a cistern and is drowned with the rest of the body unsubmerged.

The hands and the feet may be found tied together with, or without, weights attached to the cords. The indication is suicide in the event of the signs of death from drowning being present, the body however, should be carefully examined for marks of violence suggestive of foul play.

It sometimes happens that a person taking an ordinary warm bath for ablution is found dead without any external injuries. If the signs of death from drowning are present, the case will be either one of suicide or of accident, the latter being probably the result of an epileptic seizure, if they are absent, death will have occurred from syncope, in one case it was due to thrombosis of the pulmonary vein. The question of homicide in these cases arises when there is ground for suspicion as to the motives and actions of a second person. The medical evidence would be expected to prove the cause of death, which might be due to asphyxia produced in some other way than by drowning, and to give an account of the general condition of the body with regard to indications of a struggle having taken place.

Men in embarrassed pecuniary circumstances have been known to commit suicide by drowning in such a way as to suggest accident, in order that their families might be benefited by money payable at death under an insurance policy, at the present time there is no necessity for this subterfuge, for reasons explained in the chapter on life assurance. From time to time suicide is committed in this way, the suicide's object being to spare his family any reflected odium. Medical evidence can do little beyond proving the cause of death, the distinction between suicide and accident will have to be determined by circumstantial evidence.

### RESUSCITATION FROM DROWNING.

Since drowning usually results from asphyxia, with certain alterations in the physical condition of the lungs, the possibility of recovery after submersion is dependent upon two contingencies—the duration of the asphyxia, and the stage which the changes in the lung tissues have reached.

In experiments with dogs in which the trachea is suddenly obstructed, it is found that the movements of respiration continue for about four minutes, the pulsations of the heart persisting for two to three minutes longer, recovery being possible at any period short of that when the heart ceases to beat. The experiments conducted by a committee of the Royal Medical and Chirurgical



Society<sup>1</sup> show that the period within which recovery is possible is very much curtailed if drowning is the mode by which asphyxia is produced. Two dogs, one with the trachea plugged and the other not, were submerged for two minutes and then withdrawn from the water, the one in which the trachea was plugged recovered, the other did not. The reason for this difference lies in the condition of the lungs. When the trachea is plugged neither air nor water can enter them, and consequently on removal of the obstruction, they are in a relatively fit state to resume their function, when the trachea is free, water is drawn into the finer bronchi and the alveoli in the attempts at respiration, and consequently the lung tissue becomes sodden and loses its elasticity.

In human beings asphyxia supervenes in from one to two minutes after submersion, and death usually before five to six minutes, cases of recovery after longer submersion are recorded, but it is probable that in them some condition obtained which prevented water being drawn into the lungs. By practice, possibly aided by some exceptional inherent condition, individuals have acquired the capacity of enduring a longer period of submersion than ordinary without injurious results, in such cases there is obviously no attempt at breathing whilst under water, so that the question resolves itself into a capacity to postpone the asphyxial limit beyond that which is usual. In the description given of the condition of the lungs in drowning, it was stated that the aspiration of water causes a secretion of mucus in the bronchial tubes which tends to block the finer bronchi. After prolonged submersion, although the heart may continue to beat, and the vital condition (so far as the degree of asphyxia is concerned) is not hopeless, yet this blocking of the finer bronchi may prevent air reaching the alveoli after removal of the body from the water, so that attempts at artificial respiration merely churn the mucus and froth to and fro in the air-passages without furthering respiration.

It is very encouraging to know, however, that with proper treatment perseveringly carried out recovery not unfrequently occurs after prolonged submersion, even when the case appears hopeless. On one occasion a boatman towed the body of a man ashore, and pronounced life to be extinct, a medical man corroborated this opinion, but two more sanguine bystanders began vigorous treatment, with the result that the apparently drowned man entirely recovered from the effects of the submersion. A good deal depends on prompt treatment, unfortunately, it often happens that among those present no one has a sufficiently practical acquaintance with the method of carrying on artificial respiration as to put it into execution in an efficient manner. The absence of means of supplying artificial warmth on the spot is another cause of failure, the success which has attended the establishment in Paris of places of succour for the drowned demonstrates the difference between immediate and delayed treatment. Formerly, submersion for two minutes was considered the limit of the probability of success, at present, prompt treatment, with means of applying warmth in the form of hot baths and blankets always to hand, enlarges the limit to five minutes, at which recovery is said to be certain.

The following remarkable instance of recovery after prolonged submersion is recorded by Pope<sup>2</sup>—A man was sailing in a boat when it capsized and he fell into the water with some weights on the top of him, so that with the exception of his left arm he was entirely and continuously submerged for from twelve to fifteen minutes, he was resuscitated with considerable difficulty and eventually recovered. The favourable issue was attributed to the weights pressing

<sup>1</sup> *Med. Chirurg. Trans.*, 1862

<sup>2</sup> *The Lancet*, 1881

on the chest, which, together with concussion, so interfered with respiration as to prevent any water from entering the lungs

## ARTIFICIAL RESPIRATION

**Silvester's Method** —After emptying the mouth and throat by turning the patient face downwards for a few seconds, place him on his back with the head a little higher than the feet, the head and shoulders being supported on a compactly folded article of clothing, remove everything tight about the chest and neck, draw the tongue forwards and maintain it in that position. Grasp the arms just above the elbows and draw them steadily above the head, keeping them on the stretch for two seconds, then reverse the movement, and press the arms firmly downwards against the sides of the chest for two seconds, repeat these manœuvres about fifteen times every minute until natural breathing is initiated, or as long as there is any hope of resuscitation. In the meantime friction and warmth are to be applied to the body.

**Howard's Method** —Turn the patient face downwards with a firm roll of clothing under the stomach and chest, press with the whole weight on the patient's back two or three times for about four seconds each time, so as to drain the mouth and the air passages as much as possible of the accumulated water. Then turn the patient face upwards, so that the roll of clothing is just below the shoulder blades, the head hanging back. Place the patient's hands above his head, kneel across his hips, fix your elbows against your hips, grasp the lower part of the patient's chest, and squeeze the sides together, pressing gradually forwards with all your weight for about three seconds until your mouth is nearly over that of the patient, and with a push suddenly jerk yourself backwards. Pause for three seconds, and then repeat the process, which should be performed about eight or ten times a minute.

**Laborde's Method** —Turn the patient face downwards for a few seconds so as to empty the mouth and throat, then place him on the back with the head low and insert a piece of wood, or similar substance between the molar teeth so as to keep the mouth open. With the corner of a pocket handkerchief wrapped round the finger free the throat of mucus, then cover the right forefinger and thumb with the handkerchief and seize the tongue as far back as possible, pull it sharply out and immediately relax the tension, the entire tongue being thus drawn forward about fifteen times a minute. When additional assistance is available, bilateral compression of the chest should be effected synchronously with the traction on the tongue.

**Schafer's Method** —Place the patient face downwards on the ground with a folded coat under the lower part of the chest. Kneel either across or by the side of the patient, facing his head. Place your hands flat over the back, on the lowest ribs, one on each side, and gradually throw the weight of your body forward, so as to produce firm but not violent pressure on the patient's chest. By this means the air, and water if there is any, are driven out of the patient's lungs. Then raise your body slowly, so as to remove the pressure, leaving the hands in position. Repeat this backward and forward movement every four or five seconds.

The advantages claimed for this method are —The extreme simplicity of the operation and the ease with which it may be performed. The readiness with which water and mucus are expelled from the air passages. The impossibility of the air passages being blocked by the tongue falling back into the pharynx, and the efficiency of the lung aeration that is produced.

See discussion at a meeting of the Royal Medical and Chirurgical Society, December 8, 1903, when Dr Bowles stated that he had used almost exactly the method above described for the last forty five years.

It occasionally happens that, although automatic respiration is established, the patient succumbs shortly afterwards, or remains in a moribund condition for several hours, and then dies without having regained consciousness. In other cases, a temporary recovery of a more complete character takes place which may last for twenty four or more hours, followed by death, apparently from exhaustion. The treatment subsequent to re-establishment of respiration will depend on the symptoms manifested, in some cases venesection may be indicated to relieve the heart, in others stimulants and external warmth will be more efficacious.

## CHAPTER XX

**DEATH FROM ELECTRICITY AND FROM EXTREMES  
OF TEMPERATURE.****DEATH FROM LIGHTNING.**

ATTEMPTS have been made to classify the various modes in which lightning causes death, but beyond dividing them into those in which more or less severe mechanical injuries and burns are produced, and those in which death occurs without any trace of injury, nothing is gained, nor is there much profit in speculating as to the relative conductivity of the different tissues and fluids of the body. When the 'current' of a voltaic battery or of a dynamo is in question, Ohm's law holds good, with lightning it does not. The enormous difference in potential between cloud and cloud, or cloud and earth, overrides the law of conductivity as applied to voltaic electricity experimentally produced. Unlike a voltaic current, lightning does not invariably take the path of the least resistance, it will forsake a relatively good conductor and spring across a stratum of air, which offers a resistance many thousand times greater. This occasions no surprise when the enormous potential—which has been calculated at 3,601,000 volts for a flash of lightning a mile long—is taken into consideration. Whatever is the nature of the molecular changes which take place in a conductor along which a "current" of electricity is passing, it is obvious that time is involved. The duration of a flash of lightning has been computed at the twenty-thousandth part of a second, it is easy, therefore, to understand that the molecules of a substance which serves as a pathway to such a discharge must be placed under an excessive strain, which will for the moment vastly increase the resistance, the molecules are, as it were, taken unawares, and before they can adjust themselves to the stress, the resistance is probably augmented to that of the lateral path across which the lightning leaps. The erratic course frequently taken by lightning may be thus accounted for, both as regards the human body and also in respect to articles of furniture in rooms.

When a great difference in potential exists between the earth and a cloud above it, the intervening stratum of the atmosphere acts as a dielectric, the whole forming an inductive circuit, the tension increases until the resistance of the dielectric (the air) is overcome and then a discharge takes place. Any object raised above the earth's surface diminishes the resistance of the dielectric at that point to the extent to which it projects beyond the ground-level, and will anticipate the occurrence and determine the direction of the discharge, which would have been delayed until a still higher tension had accumulated if the ground-level had been uniform. If a large area of the earth's surface is free from inanimate projections—as is the case in a plain devoid of trees or buildings—the body of a man in the erect posture may determine the line of discharge.

Although a flash of lightning appears to the eye as a mere line, and is in reality only a spark passing with immense velocity from point to point, its

influence is not limited to its apparent path. What is visible is, so to speak, but the core of the discharge—where its influence is at the maximum, on every side of this line is a wide track, within the limits of which damage may be done, hence, during a thunderstorm the proximity to a tree is dangerous—the tree determines the occurrence and receives the central part of the discharge, and the immediately surrounding objects receive portions of its encircling zone.

The thermic power of a flash of lightning is very intense, but is very limited in duration, so that burns caused by lightning are frequently extensive without being deep. Although burns of considerable depth may occur on parts of the body with which the clothing is in contact, they are usually limited to those parts underlying metallic objects—as buckles and bracelets. The sudden high temperature acquired by metallic bodies surrounded by relatively imperfect conductors is probably due to their acting as condensers, and also to their molecular constitution being of such a nature as readily to transform the electricity they receive into heat which is often extremely localised, of this Key<sup>1</sup> gives a good example. A watch and a match-box both of silver, were found together in the waistcoat pocket of a man who had been killed by lightning—although a hole a quarter of an inch in diameter had been fused through the outer case of the watch, the match-box contained two unignited wax matches. Burns may also occur on the unclothed parts of the body, as, for example, the face. Clothing saturated with rain will to a limited extent act as a conducting envelope, and will tend to divert a portion of the discharge and to lessen the amount of mischief which might otherwise be sustained by the body.

The clothing, whether dry or wet, is frequently torn, in some cases to shreds, and stripped off the body. One or both of the boots may be forced off the feet, or the soles may be stripped away from the upper leathers, sometimes a hole is simply punched through part of the boot. This mechanical effect is due to the molecular disturbance which invariably attends an electric discharge. When a discharge passes along a good homogeneous conductor, such as a piece of metal, if it is connected with the earth and is of sufficient sectional dimension, no obvious change is produced, if such a conductor, not in connection with the earth, receives a powerful discharge, it will either be damaged itself, or objects between its lower end and the earth will be damaged. A solid dielectric so interposed is either pierced or shattered when the potential exceeds the inductive capacity, a sheet of thick glass, for example can be perforated by a strong discharge from a Leyden battery concentrated upon a limited area. When the conductor is composed of a number of varying resistances, the sum of which is relatively great, it will be torn or injured by the passage of a discharge at a potential like that of lightning. This results from unequal strain, which to some extent resembles that caused by the intercalation of a series of dielectrics in the path of a discharge, at each of which it becomes disruptive, mechanically breaking them down *seriatim*, and tearing apart the structures or fabrics along which it passes. After doing a certain amount of damage the discharge may leap a distance, in the intervening space there will be absence of mechanical injury.

It is possible that the intense heat developed along the path of the discharge may also cause mechanical violence by suddenly converting water, or fluid into the composition of which water enters, into steam, in this case the chief factor would be the extreme rapidity with which the liquid is made to acquire the gaseous form.

<sup>1</sup> *The Lancet*, 1895

A peculiar appearance due to arborescent markings is not unfrequently observed on the surface of the bodies of those who have been struck by lightning. These marks are not determined by the course of capillary vessels or other anatomical structures, but are caused by divarications of portions of the discharge producing a kind of specialised erythema, which indicates the paths taken by the discharge, they resemble, both in appearance and in causation, the well-known Lichtenberg figures of experimental electricity. Distinction must be drawn between them and marks due to superficial burns, which occasionally present peculiar outlines—the latter sometimes take the form of metallic objects worn on the body, which, heated by the discharge, produce burns of corresponding shapes. Fig 18, taken from the *Lancet* (1883), gives a good idea of the arborescent markings produced by lightning.

In those cases of death from lightning in which no sign of injury is found, neither on the surface of the body nor on the clothing, it is possible that the result may be due to the victim having been just outside the path of the main discharge, but sufficiently near as to receive a shock violent enough to permanently paralyse the central organs of the nervous system. In some such cases internal examination fails to demonstrate the proximate cause of death, in others there are indications of cerebral hæmorrhage, or of diffuse disorganisation of the brain structure.

Hennessy<sup>1</sup> relates the case of a man who was killed by lightning without there being any trace of burning or injury to body or clothing, but on removing the calvarium, about half a pint of blood escaped from below the dura, although the brain itself appeared healthy. In contrast to this case is one related by Wilks.<sup>2</sup> A man struck by lightning whilst in the act of urinating in the open air, was thrown to the ground, but was not rendered unconscious. Although completely clothed before being struck, when found a few minutes after he was naked, with absolutely nothing on except part of the left arm of his flannel vest, the field around was strewn with fragments of clothing. The clothes, which were wet with rain, were split from top to bottom, and the edges torn into fringe-like shreds, they only showed marks of burning where they had been in contact with metallic objects. The hair on the face was burnt, and the body was covered with burns, which were superficial on the chest, but deeper on the abdomen and the right thigh. The right tibia was fractured—the ends of the bones projecting through the skin, and there was a lacerated wound on the right heel, with a comminuted fracture of the os calcis. The man's watch had a hole fused through it, and the chain was almost entirely destroyed, only a few partially melted links remained. A highly interesting statement was made by the patient to the effect that when urinating he habitually lifted the right heel from the ground—had it rested on the wet earth it would probably have escaped. Complete recovery took place. In a case recorded by Whichello,<sup>3</sup> two men driving in a dogcart were simultaneously killed by lightning. The body of one man was uninjured, that of the other showed a number of superficial circular burns, from a sixteenth to a quarter of an inch in diameter, on the chest and abdomen. The collar stud was fused and the skin beneath deeply burnt. The vest and shirt were charred, but the

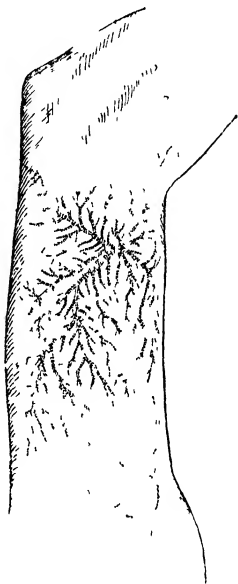


Fig 18

<sup>1</sup> *Brit. Med. Journ.*, 1889

<sup>2</sup> *Trans. of the Clinical Soc. Lond.*, 1880

<sup>3</sup> *The Lancet*, 1899

waistcoat and coat were uninjured. From the neck to the buttocks less severe burns were present, and the drawers and trousers had a hole burnt in them corresponding to the burn on the right buttock. The cushion on which this man sat was superficially burnt.

**The post-mortem appearances of death from lightning.** **External.**—Cadaveric rigidity not unfrequently occurs at the usual period, but sometimes it comes on immediately after death, in which case it is evanescent. The pupils may be dilated, contracted, or unequal. Arborescent markings, burns, singeing of the hair, ecchymoses, lacerations, or other indications of injury, may or may not be present. The clothing may be intact or it may be torn and stripped off the body, it may show signs of burning. Partial fusion of coins or other metallic objects on the person of the deceased is very significant of the mode of death. Kratter<sup>1</sup> directs attention to this means of diagnosis when other indications are wanting. Steel implements, such as pocket-knives, are often magnetised.

The **internal** signs are not characteristic. The blood has been described as dark and fluid, it is so sometimes, but it may coagulate as usual after death. The membranes and vessels of the brain have been found hyperæmic, and, in some cases, blood was extravasated. Fractures of the bones, including those of the skull, have occurred, but only exceptionally. Putrefaction usually commences early.

On the average about sixteen persons are killed by lightning annually in England and Wales. The death of males largely outnumber those of females owing to much greater outdoor occupation.

When an individual who is struck by lightning escapes death, it is not unusual for anomalous nerve symptoms to manifest themselves, the symptoms may take the form of sensory or of motor disturbances, with or without derangement of the special senses. In the case related by Paige, Buller, and Mills,<sup>2</sup> a woman was rendered insensible by being struck by lightning which had apparently touched the left side of the forehead above the eyebrow. A few minutes after she was found motionless, unconscious, and in a state of muscular relaxation with the left eye closed, the right eye open, the face purple, and the pulse imperceptible, neither heart nor respiratory sounds were audible. On recovery of consciousness, it was found that the pharyngeal muscles were paralysed, as well as those of the upper part of the body, including the arms, the right eye could not be moved, the patient could hear, but could not speak. It was two weeks before the pharyngeal paralysis had passed off sufficiently to allow solid food to be swallowed. For five or six weeks the left eye was turned inwards and upwards, occasioning diplopia, ophthalmoscopic examination showed pallor of the optic nerve, but nothing indicative of neuritis past nor present, the field of vision was uniformly restricted for all colours. All the symptoms eventually disappeared, except that the left eye remained weak.

Sometimes the spinal cord is rendered functionless for a time, producing paraplegia, with anæsthesia of the paralysed parts, the bladder sharing in the paralysis, in other cases, the psychical powers are disordered, the patient becoming intellectually enfeebled, or melancholic. A common condition is that of timorousness, especially during thunderstorms, which is probably simply the result of psychical shock, such as is caused by other alarming occurrences.

When examining a body found under circumstances which render death from lightning probable, the possibility of it having resulted from other causes must not be overlooked. Schauenburg<sup>3</sup> states that the dead body of a woman was found in a wood at first it was thought that she had been struck by lightning, but examination showed that death was due to accidental strangulation caused by the bonnet-strings being drawn tightly round the throat in consequence of a fall. Hofmann<sup>4</sup> relates the case of a girl who was in a room

<sup>1</sup> *Vierteiljahrsschr. / ger. Med.*, 1891

<sup>2</sup> *Casper's Vierteiljahrsschr.* 1855

during a violent storm which blew in the casement of a window. The shock broke a pane of glass, and a fragment the shape of a knife-blade penetrated her chest, and caused death from internal hæmorrhage, although no one had heard any thunder, the sudden death was attributed to lightning, and the body was buried, it was not until three weeks after that the actual cause of death was explained.

### DEATH FROM THE ELECTRIC CURRENT.

Death from the **electric current** occurs from time to time owing to the body coming into accidental contact with conductors conveying direct or alternating currents of high tension. It is commonly believed that the alternating current is more dangerous to life than the direct current, probably because fatal accidents occur more frequently with the former than with the latter. This, however, is due to the tension rather than to the type of the current, alternating currents are usually generated at high tensions, hence the danger, at equal tensions the direct current, to say the least of it, would be quite as deadly as the alternating current. Whatever may be the pressure in the street-mains, the regulations of the Board of Trade (except with its express approval and only for special purposes) forbid the introduction into dwelling-houses of currents exceeding 250 volts, therefore, when the main current is of high tension—1,000 to 3,000 volts—transformers, by which the necessary reduction in pressure is effected, are interposed between the main and the house supply. A pressure of 225 volts has proved fatal, if the pressure is much below this there would be little or no danger to life, in addition to the pressure, however, the duration and mode of contact have to be taken into account. The effect of good contact and consequently of low resistance is shown in the case of a man who, when standing barefoot on a mixture of potash and sugar, was killed by a current of 95 volts. In another case a tension of 65 volts caused death. A man, in order to avert falling from a ladder, seized with both hands the two non-insulated leads of an electric-light circuit with a potential of 240 volts and was immediately killed.<sup>1</sup> On the other hand, recovery has taken place after the passage of 2,500 volts, and in another case after 5,500 volts. Fatal discharges may be received without the body being actually intercalated in the main circuit, as, for example, when the current passes from one of the conductors through the body to earth.

Two views are held as to the way in which the electric current causes death. One is that it paralyses the heart, the other that it arrests respiration. Experiments on animals have so far yielded discordant results. Bleile, in a paper read before the American Institute of Electrical Engineers, 1895, states that arrest of the heart beat and not failure of respiration is the cause of death. This opinion is shared by Tatum.<sup>2</sup> Oliver and Bolam<sup>3</sup> as the result of experiments with the alternating current on dogs and rabbits, state that with a potential sufficiently high to cause death, but not excessively high, death is due to heart paralysis, the breathing being rhythmically continued for a short time after the heart has ceased to beat, when the potential is considerably higher than that usually required to kill the animal, simultaneous stoppage of the heart and lungs occurs, but under no circumstances is failure of the heart preceded by arrest of respiration. On the other hand, d'Arsonval<sup>4</sup> believes that death is usually due to arrest of respiration, this view is supported by Kratter,<sup>5</sup> whose experiments lead him to the conclusion that death from electricity is nothing but a special kind of suffocation due to primary stimulation and subsequent paralysis of the respiratory centres. Both of these observers admit that exceptionally death may be due to heart paralysis. Jellinek<sup>6</sup> states, from experiments on animals, and from

<sup>1</sup> *Nature*, 1900

<sup>2</sup> *New York Med Journ*, 1890

<sup>3</sup> *Brit. Med. Journ*, 1898

<sup>4</sup> *Arch. d'Anthropol. Crim*, 1895

<sup>5</sup> *Mittheilungen d. Aerzte in Steiermark*, 1895

<sup>6</sup> *Wiener Flin. Wochenschr*, 1905

experience in the case of human beings, that there is no specific mode of death from electricity. psychical effects of the nature of shock are associated with certain dynamic effects on tissues which may be simply inhibitory, involving no histological changes, or may cause obvious alternation of structures. Jex Blake has re-examined the whole subject<sup>1</sup>

In human beings the symptoms produced by electric discharges comprise — Local injuries, stertorous breathing, insensibility, pallor of the face, or congestion with suffusion of the eyes, cold, moist surface, and dilated pupils, in more severe cases there is immediate insensibility, with perhaps one or two gasps, and death. The post-mortem appearances include burns or marks on the parts of the body which came in contact with the conductor, hyperæmia of the internal organs, œdema of the lungs, fluid blood, and ecchymoses, particularly along the path taken by the current. According to Kratter, the post-mortem appearances are those of asphyxia, the heart, however, has been found empty.

With alternating currents of very high pressure peculiar local injuries may be caused, in one instance a workman momentarily touched a live wire in such a way as to receive the current through one hand and arm only, the end of the thumb which touched the wire was split open as though by an explosive. Shield and Delépine<sup>2</sup> examined the skin in the region of a blister on the finger of a man who was accidentally killed by touching a dynamo, and gave a detailed account of the microscopic appearances, which differed from those of ordinary burns.

**Treatment.**— The first thing is to switch off the current, or, if that cannot be done, to detach the person from the conductors if he is not already free, the rescuer must on no account use his bare hands, but (in the absence of rubber-gloves) should protect them with non-conducting material, such as several folds of dry cloth. The sufferer may be drawn away by his coat *if dry*, or a convenient plan is for the rescuer to take off his own coat and thrust his hands into the sleeves, he can then withdraw the body with little or no risk. Artificial respiration should be resorted to at once and persevered in for at least two hours if necessary. It is said that the great majority of cases recover if first-aid is promptly rendered. Venesection is advised by Kratter.

### DEATH FROM HEAT-STROKE.

This cause of death is also called **sunstroke**; but since death frequently occurs without the direct action of the sun's rays, the latter term is too restrictive. Heat-stroke is probably due to derangement of the heat-regulating centres by continued high temperature of the surrounding air, particularly when associated with excessive humidity, under circumstances which place the individual attacked in an unfavourable condition for resistance. It mostly occurs in tropical countries, especially where large bodies of men are massed together (as in barracks). Soldiers on march, or civilians taking an unwonted amount of exercise in the open air when the rays of the sun are excessively powerful, are liable to be attacked, either suddenly, whilst the exercise is being taken, or in a few hours after, when the weather is excessively hot, patients are not unfrequently attacked during the night.

**Symptoms** — Occasionally, the onset is abrupt, but more frequently there are prodromal symptoms of headache, drowsiness, and nausea. A frequent desire to micturate is mentioned by some writers as an early symptom. The temperature rises suddenly to 104° F or more, and in severe cases it may reach 110° F.

<sup>1</sup> *Brit Med Journ*, 1913

<sup>2</sup> *Brit Med Journ*, 1885



The skin is hot and dry, and occasionally shows a macular eruption. Delirium may occur, but coma is more frequent. The pulse is rapid, and becomes irregular, intermittent, and weak. The motions may be passed involuntarily. Respiration is noisy and quick. In fatal cases convulsions occur, the respirations become slower and more stertorous, and the patient dies of asphyxia in from a few hours to a day or two. In favourable cases the temperature falls rapidly, but the patient remains exceptionally susceptible to high temperatures for a long time. Castellani and Chalmers state that the death rate varies between 15 and 25 per cent, but it may be as high as 51 per cent.

The **post-mortem appearances** are—early onset and disappearance of cadaveric rigidity, with rapid advance of putrefaction, hyperæmia of many of the internal organs, such as the brain (which may be œdematous), the lungs, and the vessels of the splanchnic area. Microscopically, the minute vessels, especially of the medulla, are very congested, and the nerve cells show coagulative necrosis and disappearance of the Nissl bodies, together with a swollen and chromatolytic condition of the nucleus, though the nucleolus may remain apparently intact (Castellani and Chalmers).

**Heat syncope** or **heat exhaustion** is a condition occurring among persons whose bodily health is abnormal when they are subjected to high temperature and excessive humidity of the atmosphere. It is particularly apt to occur in those suffering from chronic alcoholism with fatty infiltration of the heart muscle. The symptoms come on suddenly with giddiness, nausea or vomiting, and staggering gait. The pulse is small, the skin pale and cold, and the temperature subnormal. Most cases recover, but at times coma supervenes followed by death.

In 1919, the deaths of 63 males and 35 females were attributed to heat-stroke in England and Wales.

## DEATH FROM COLD.

Exposure to extreme cold in the first instance causes contraction of the vessels of the skin, the blood being driven into the internal organs, subsequently, if the exposure continues, the superficial blood-vessels are paralysed, giving rise to patches of localised erythema.

The **symptoms** are loss of energy, both physical and mental, followed by drowsiness and disinclination to move. The mental faculties become torpid, the special senses numbed, and the victim is seized with an irresistible desire to lie down and sleep, if this is yielded to, and help is not forthcoming, the lethargy passes into profound sleep, which deepens into stupor, and finally death takes place from general depression of the systemic powers. In some cases special disturbances of the nervous system manifest themselves in the form of clonic spasms or of physical derangement.

It is probable that death results primarily from diminished supply of oxygen to the various organs, especially to the nerve-centres. A very low temperature interferes with the capacity of hæmoglobin to give up its oxygen. Pflüger<sup>1</sup> found that at 0° C. oxygen cannot be liberated from combination with hæmoglobin by removing the atmospheric pressure with the air-pump, as Bunge<sup>2</sup> puts it, the dissociation of oxygen from hæmoglobin is due to heat and not to the vacuum. At a low temperature the ordinary reducing agents, such as ammonium sulphide, take a much longer time than usual to abstract oxygen from hæmoglobin. Given, then, a diminished capacity of the hæmoglobin

<sup>1</sup> *Arch. f. gesammte Physiologie*, Bd. 1.

<sup>2</sup> *Lehrbuch d. phys. u. path. Chemie*, 1889.

to yield up its oxygen, and a coincident lowering of the activity of the tissues as regards their power to utilise it (internal respiration), an explanation is afforded of the gradual and general depression of the systemic powers and, also, of the occasional manifestations of disturbance of the nerve-centres.

In addition to these essential conditions certain influences of external relation frequently come into play. Preliminary fatigue, with or without deprivation of food and sufficient clothing, renders the victim more susceptible to the effects of exposure to cold, recourse to alcohol in order to stimulate the jaded powers increases the desire for sleep, the extremes of age and feeble health diminish the capacity of resistance to cold.

The **post-mortem appearances** of death from exposure to cold, with one exception, are not characteristic. The appearance alluded to is that produced by cherry-red coloured spots or stains on those parts of the surface of the body which are exposed to the air, they strongly resemble in colour the stains met with in cases of poisoning by carbon monoxide. Rollet<sup>1</sup> explains the formation of these stains on the assumption that, in consequence of the extremely low temperature, the red corpuscles part with their hæmoglobin, which dissolves in the plasma, forming 'lake-coloured' blood, thus, exuding through the walls of the vessels, tinges the neighbouring tissues red. To some extent this may be the case, but Falk<sup>2</sup> gives the most satisfactory explanation of the way in which the stains are produced. When hæmoglobin is exposed to an extremely low temperature it loses its property of parting with oxygen, but not that of combining with it, this applies to the blood both in the living and in the dead. The consequence is that when a body is exposed to a temperature below the freezing point, the blood in the superficial vessels of those parts which are exposed to the air acquires oxygen by diffusion through the skin, and, retaining it, assumes the bright-red hue of arterial blood, within certain limits, if the temperature remains at or below zero, the cherry-red colour persists. When a body which displays this appearance is exposed to a higher temperature the hæmoglobin, regaining its capacity of yielding up oxygen, is reduced by the oxidising properties of the surrounding tissues which now come into play—the stains consequently darken and become like those after death from ordinary causes.

The cherry-red coloured stains on those parts of the body exposed to air at a low temperature is not a proof of death from cold, a similar appearance may be produced after death from other causes if the body is subsequently thus exposed. It is only on the parts of the body exposed to the air that the blood assumes this peculiar colour, the parts which are covered, either by the clothing or by contact with the surface on which the body rests, present the usual appearance.

The **internal appearances** are limited to hyperæmia of the vessels of the brain and its membranes, with occasionally a similar condition of those of the abdominal viscera. Wichniewski<sup>3</sup> examined forty-four cases of death from extreme cold, and invariably found under the epithelium of the gastric mucous membrane small extravasations of blood which can be lifted off with the point of the scalpel without visible loss of substance, the rest of the gastric mucous membrane is hyperæmic, thickened, and corrugated. On account of the access of oxygen, the lungs are often bright in colour, like that of the exposed post-mortem stains on the surface of the body. The heart may be distended with blood, which is usually dark, but has been found bright in colour. From experimental

<sup>1</sup> *Sitzungsberichte d. Kaiserlichen Akad. d. Wissenschaft*, Bd. xlv.

<sup>2</sup> *Vierteljahrsschr. f. ger. Med.*, 1887 u. 1880.

<sup>3</sup> *Arch. de l'Anthrop. Crim.*, 1895.

researches on animals, Falk is disposed to regard a bright-red hue of the blood in the heart as a sign of death from cold. It is not produced by exposing dead bodies to the influence of a low temperature, because the atmospheric oxygen cannot diffuse itself so far internally as the heart. In animals that have died from exposure to a very low temperature which is continued after death, the blood in the heart acquires the same hue as that in the lungs and on the surface, hence, whilst dark-coloured blood in the heart is consistent with death from cold, bright-coloured is indicative of it.

If putrefactive changes have commenced before the body is exposed to a low temperature, the characteristic stains do not appear, because tissues undergoing decomposition are active deoxidisers, even at the freezing-point, decomposing tissues take up all the oxygen that is diffused through the skin to the neglect of the hæmoglobin, which thus remains in the condition of reduced hæmoglobin. From this it appears that although putrefaction in the broad sense is arrested at and below the freezing-point, a slow process of oxidation goes on in the parts which are accessible to air *in cadavers in which decomposition has already set in*.

When an individual dies from exposure to a low temperature (at or below the freezing-point), to which the body has been continuously exposed after death, the changes due to putrefaction will be absent. The same absence of putrefaction occurs when death has resulted from other causes than cold, if the dead body is immediately and continuously subjected to a temperature below 0° C. Under these conditions bodies have been preserved without apparent change for an indefinite time. An interesting account is given in d'Arve's *Histoire du Mont Blanc* of the discovery of the remains of some guides who were swept down a crevasse by an avalanche in the year 1820, where they remained buried in ice at a great depth. In the year 1861, after an interval of forty-one years, they were discovered. Amongst the rest, a forearm and hand with the fingers intact were found, the flesh was white and fresh looking, and the finger-nails were rose-red when first seen, but the colour rapidly faded, on the ring-finger was a slight abrasion, the blood-stain from which was quite visible. The joints of the arm and of the hand were flexible when thawed, as is the case in a body recently dead.

If a dead body is found in the frozen condition with signs of putrefaction present, the assumption is that death was not caused by cold, there exists, of course, the remote possibility of an elevation of the surrounding temperature after death—during which putrefactive changes might have taken place—followed by renewed cold.

A frozen dead body is perfectly rigid, like a marble statue. When the body is thawed, it is not unfrequently found that cadaveric rigidity is present, although the period after death when it usually ceases has long since passed, this applies to the bodies of those who have died from the effects of cold, and also of those who have died from other causes if they have been immediately subjected to a temperature below 0° C. After a frozen cadaver has been thawed, putrefaction quickly commences and advances very rapidly.

In 1919, the deaths of 54 males and 17 females were attributed to exposure to cold, in England and Wales.

## CHAPTER XXI

## STARVATION.

**Starvation** may be caused either by absolute deprivation of food or by insufficient amount or defective quality. Cases which come under the notice of the medical jurist are usually those of young children. If adults, they are generally persons of feeble intellect who have been isolated in a cellar or some other unfrequented part of the house and entirely neglected both as regards nourishment and bodily attention. This treatment is adopted by those who put it in force with the design of ridding themselves of the presence of the victim without daring to kill him in a more direct manner. From parsimonious motives, servant girls have been deprived by their employers of sufficient food to support life. If the person undergoing starvation is rescued alive, medical evidence will be required to substantiate the charge of criminal neglect from the appearances present in the living subject.

The degree of emaciation after deprivation of food for a given period depends to a great extent upon the bodily condition at the time the starvation was commenced. For this reason, in cases of fatal chronic starvation in which the minimum amount of food to sustain life has been given for a prolonged period, followed by an interval of total deprivation, the degree of emaciation is much greater than when a well-nourished healthy individual is suddenly cut off from food until he dies.

**Symptoms.** The natural feeling of hunger which indicates to a healthy person that the organism is in want of food, disappears after the first thirty-six or forty-eight hours of fasting, there may be pain and discomfort in the region of the stomach but it is not associated with a desire for food. Intense thirst is always present, and want of fluid greatly increases the sufferings. Muscular weakness gradually occurs, and is quickly attended by emaciation, which is progressive until the end. The skin is wrinkled and is usually pale, dry, and of a parchmenty appearance. purpuric spots have been noticed, and, more frequently, brownish stains resembling ingrained dirt. The features and eyes are sunken, and the malar bones stand prominently out. The mouth and tongue are dry, and the breath has a disagreeable odour, which in some stages of starvation is of an ether-like nature, at a later stage the whole body gives of a peculiar putrescent odour, but unlike that of ordinary putrefaction. The abdominal wall is concave, so that the bodies of the vertebræ may easily be felt through it, and to use the popular phrase, the limbs are little more than skin and bone. The mind may remain clear or may be enfeebled to imbecility, hallucinations are not uncommon. The **pulse** is feeble, but does not present any constant numerical characteristic, it is often slower, and sometimes quicker, than in the normal condition. Two distinct statements are made with regard to the **temperature**: one is, that it does not fall until shortly before death, when it quickly diminishes, and the other is that the temperature is subnormal throughout. Some of this discrepancy may be due to a blending of observations made on man and on animals. Falk,<sup>1</sup> from experiments on animals, states that

<sup>1</sup> *Beitrage z. Physiol*, 1875

the temperature remained constant for fifty-three days, and then rapidly went down till death took place. Paton and Stockman,<sup>1</sup> when investigating the case of a man who fasted thirty days, found that the temperature ranged from 96° to 93.4° F. The amount of oxygen consumed by starving animals is not greatly below the normal, and, therefore, the temperature may be maintained at, or a little under, the normal point.

The **loss of weight** varies with the state of nutrition which exists at the commencement of starvation. A stout, well-nourished man, cut off from food, will lose very much more weight during the earlier days than a spare man; after three or four days' fasting, their ratios become more equal. Paton and Stockman found that a man in a thirty days' fast lost 10.316 grms, or 166 kilo per kilo of original weight, in all, about one-sixth of the original weight, the daily loss averaged 31 kilo. Falk<sup>2</sup> states that warm-blooded animals lose about 40 per cent. of their original weight before they die from total deprivation from food and drink.

The daily amount of urine is lessened and, when no fluids are taken, the diminution is progressive, the colour and specific gravity being higher than normal. The daily excretion of urea during starvation has been estimated by countless observers. When food is cut off, the excretion of nitrogen (urea) suddenly drops and then remains fairly constant for several days with a slight tendency to further diminution. When all the fat is used up, there may even be a little increase, which, in any case, is only temporary, and gives place to a progressive fall until death takes place. Luciani,<sup>3</sup> in his observations on Succi during one of his fasts, found that on the last day of taking food the N in the urine amounted to 16.29 grms. During the first twelve days of fasting it sank from 13.8 grms to 7.29 grms and progressively diminished to 3.2 grms on the twenty-second day. Water was taken in small amount all the time, the daily amount of urine varied from 600 to 250 c.c. Paton and Stockman give the following daily average amounts of N for six consecutive periods (each consisting of five days) during a fast of thirty days: (1) 11.9 grms, (2) 5.4 grms, (3) 5.1 grms, (4) 4.2 grms, (5) 4.2 grms, (6) 3.1 grms. After the sixteenth day water was taken *ad lib*. Of the other constituents of urine—uric acid, sulphuric acid (in all its combinations), and phosphoric acid progressively diminish during starvation. The excretion of chlorine is not so constant, in some cases it has been observed to diminish progressively, and in others, at the end of prolonged deprivation from food, to very nearly equal in daily amount that given off at the commencement. Kreatinin is another irregular constituent, it diminishes during the earlier period and may subsequently increase. Baldi<sup>4</sup> found it present in very small amount during fasting. The urine may contain sugar (starvation glycosuria) diacetic acid, and acetone.

The daily excretion of CO<sub>2</sub> is diminished, but not so rapidly as the excretion of urea. In the earlier stages, especially if the animal subjected to starvation carries much fat, a considerable diminution in body weight takes place before the output of CO<sub>2</sub> is much lessened, afterwards it progressively diminishes.

The secretion of **gastric juice** during starvation varies. By passing a tube into the stomach Pock<sup>5</sup> sometimes obtained only a few drops, and at others 28 c.c. or more, the juice thus obtained often contained no HCl. If to the juice devoid of HCl a little of the acid was added it behaved as to power of digestion like normal gastric juice. Schreiber<sup>6</sup> states that the gastric juice continues to be secreted during fasting, and that, for the most part, it contains HCl, other observers have found HCl frequently absent. The daily amount of **bile** diminishes as starvation progresses, but the gall bladder is usually found distended after death. The **fæces** are at once diminished, and very soon are entirely absent.

Groll<sup>7</sup> found that the hæmoglobin is less quickly used up than the other solid constituents of the blood. Raum<sup>8</sup> found the blood colouring matter in dogs increased during hunger.

**Post-mortem Appearances.**—Cadaveric rigidity may be present, as after other modes of death. The skin is dry, with scurfy surface, or it may be horny.

<sup>1</sup> *Proc. of Roy. Soc. Ed.*, vol. xvi.

<sup>2</sup> Maschka's *Handbuch*.

<sup>3</sup> *Fisiologia del digiuno*, 1889.

<sup>4</sup> *Lo Sperimentale*, 1890.

<sup>5</sup> *Prager med. Wochenschr.*, 1889.

<sup>6</sup> *Arch. f. exper. Pathol.*, 1888.

<sup>7</sup> *Pflüger's Arch.*, 1888.

<sup>8</sup> *Arch. f. exper. Pathol.*, 1891.

and, on account of absorption of the fat, is with difficulty separated from the underlying muscles, in chronic starvation fat is almost entirely absent throughout the body. The muscles are atrophied and soft. The brain and cord show no appreciable changes. The lungs are paler and contain less blood than usual. The heart is small and may contain blood in all its chambers. The whole of the intestinal tract is atrophied. The stomach is contracted, so as often to appear like part of the large intestine, and it may contain a little bile stained fluid, the mucous membrane, usually corrugated and pale, may be slightly reddened. The small intestine is contracted, and is empty or contains a small quantity of bile. The large intestine may contain a trace of hard faecal matter. The rectum may be hyperæmic. The walls of the intestines, especially of the duodenum, jejunum, and ileum, are much atrophied, and in consequence are almost transparent, sometimes so much so as to render any substance within them easily visible. The liver is reduced in size, its surface being smooth and its colour dark. The gall-bladder is almost invariably distended with bile. The pancreas is atrophied, often to disappearance. The urinary bladder is contracted, and is usually empty or nearly so. The blood contains less than the normal amount of water.

The **appearances** which are **most characteristic** of death from starvation are—disappearance of the body-fat, and general atrophy of the soft structures. The disuse of the digestive tract results in well-marked and distinctive atrophy of its walls, and absence of stimulation to the flow of bile causes it to accumulate in the gall-bladder. The **order** in which the **tissues waste** is—first the fat, then, the glandular organs, and afterwards, the skeletal muscles.

It will be observed that the signs of death from starvation are not such as to enable a positive opinion to be expressed as to its occurrence. Similar indications are met with in death from various wasting diseases of organic origin, and also in some neuroses which leave no post-mortem evidence of their occurrence—a most important fact and one to be remembered by the medical witness. Before giving a definite opinion that death resulted from starvation, not only must all the characteristic signs which accompany that mode of death be present, but there must also be an entire absence of all indications of disease. The pathological conditions to be specially borne in mind are—**Constriction of the œsophagus**, either from simple stricture, from malignant disease, or from external pressure, such as may be caused by an aneurism. **Malignant disease** in any part of the body. **Tubercle**, miliary and otherwise, in relation to which the condition of the pia mater, peritoneum, glands, and joints should be specially examined. **Addison's disease** is not always accompanied by emaciation, but the adrenals should always be examined. The possible occurrence of **diabetes** necessitates the examination of the urine for sugar. **Chronic diarrhœa** or dysentery are other causes of emaciation which require consideration at the time the necropsy is made. In addition to these more or less obvious causes of general atrophy, neuroses, such as hysteria, hypochondriasis, or insanity, are to be taken into consideration when an accusation is brought against a person, or persons, for having caused the death of any one under their care by withholding food.

The case of *Reg v Staunton and Rhodes* (Cent Crim Court, 1877) affords a good illustration of the circumspection demanded of the medical witness when called upon to determine the cause of death in a case of alleged criminal starvation. Harriet Staunton, aged thirty-five, the wife of the first named prisoner, was stated to have been starved to death by her husband and his paramour and co prisoner, Alice Rhodes. Shortly before her death, Harriet Staunton was removed to Penge and was there seen by a medical man, she died

directly after. A certificate of death from natural causes was given, but suspicion being subsequently aroused as to the real cause of death a necropsy was made, and the case was tried as above stated. The examination revealed great emaciation and a generally filthy, neglected condition of the body, which measured 5 feet 5½ inches and weighed 74 pounds, two and a half years previously the woman weighed 119 pounds. The brain was healthy, but there were a few milary tubercles deposited in the pia mater. The heart was small. The lungs were healthy, except for a small patch of tubercular deposit at the apex of the left lung. The gall bladder was full of bile. The coats of the stomach were thinned. The intestines were shrivelled and empty, the rectum was hyperæmic for about 4 inches, the rest of the intestines being pale in colour. There was a total absence of fat with general atrophy of all the organs. The bladder contained about three ounces of urine. Positive medical evidence was given at the trial that death was due to starvation, and the prisoners, who included Staunton's brother and his wife, were convicted and sentenced to death. The medical evidence was subsequently freely criticised, emphatic opinions being expressed that there was no medical proof of death from starvation. The line of objection taken was that no proper search had been made for other possible causes of death: the œsophagus was not examined, the urine in the bladder was not tested for sugar, and the adrenals were not examined, although there was discoloration of the skin on the face. The absence of thinning of the intestines and the presence of tubercle both in the membranes of the brain and in the lungs were held, on the one hand, to negative death from deprivation of food, and, on the other, to show the possibility of death from disease. The result was that Rhodes was pardoned and the capital sentences passed on the other three prisoners were commuted.

The lesson to be learnt from this case is valuable, and is applicable to all medico-legal investigations. A medical practitioner should never approach a case with a pre-formed opinion as to its nature, he should never neglect making a searching examination of all the organs, and he should never ignore pathological indications that do not happen to agree with an opinion that has been formed, but should rather ponder as to their import.

The time required for death to result from deprivation of food varies according to certain conditions. Age exercises a considerable influence, very young children quickly succumb, adults resist better, and old people best of all. The better the health and the nutrition of the body at the commencement of the fast, the greater the power of endurance. *Drinking water tends to prolong life considerably.* Conservation of the body-heat is another favourable factor, a person well clothed, especially if so placed that the surrounding air is still, and of a moderate temperature, will live longer without food than one who is exposed to the action of cold. Colletta<sup>1</sup> states that a girl, aged sixteen, was buried under the ruins caused by an earthquake. She remained underground without food or drink eleven days, holding in her arms an infant that died on the fourth day, when extricated, the corpse of the infant was undergoing decomposition, but, being hemmed in, the girl, who was alive when rescued, was unable to rid herself of it. Life has been prolonged fourteen days, and it is stated twenty days, without food or water. In the absence of food, but with access to water, a man has lived sixty-four days.

<sup>1</sup> *Storia del Reame di Napoli*, 1820

## CHAPTER XXII

## DEATH CAUSED BY BURNS AND SCALDS.

**Burns** are caused by the action of a temperature considerably above that of the human body which is brought to bear upon it in the form of radiant heat or flame or from contact with heated solid bodies. Injuries caused by solid bodies which become liquid at an elevated temperature—such as metals in a state of fusion—are classified as burns. Substances which act chemically and produce corrosion are also said to cause burns, injuries of this kind will be considered apart from the rest.

**Scalds** are caused by the action of steam or heated liquids on the surface or in the mouth or other cavities of the body.

The **injuries** produced are proportionate to the temperature, the dimensions, and the period of action of the causal agent. The **danger to life** depends upon the severity of the injuries produced, both as regards the depth to which the tissues are disorganised and also as to the extent of the superficial area involved, and to some degree upon the localisation of the lesions.

A severe burn or scald of small superficial area, provided that its localisation is not exceptionally dangerous, is less risky to life than one which implicates a large superficial area, but does not penetrate so deeply, if a superficial area equal to one-third of the entire body-surface is destroyed, death is almost certain to ensue. This leads to the consideration of the **cause of death** from burns and scalds.

**Death** from burning may take place **immediately** the injuries are inflicted, within a **short time** after—from a few hours to several days, or at a more **remote period** when inflammation and suppuration have set in. The cause of death during the last-mentioned stage is self-explanatory, that involved in the two preceding stages is not yet determined, it is, therefore, necessary to consider briefly the most important of the many theories that have been advanced for its elucidation.

Some of these theories are based on assumptions mutually opposed. One<sup>1</sup> is that the vessels of the skin being destroyed by the heat, the blood that under normal conditions would circulate in them is driven into the internal organs, causing hyperæmia and ecchymoses, which lead to death. Another<sup>2</sup> and contradictory interpretation is that the vessels of the skin are dilated, and that death is occasioned by the resulting excessive cooling of the body, such as is supposed to occur when the surface of the body is experimentally covered with an impervious varnish, according to this view the hyperæmia of the internal organs is accounted for by the loss of tonus of the blood vessels. Another<sup>3</sup> theory, also founded on the loss of skin function, assumes the retention of certain excretory products which cause death by auto-intoxication. These products are regarded by some to be physiological, by others, pathological—that is, abnormally constituted excretory products, or toxins, which partake of the nature of alkaloids—*i.e.*, derivatives of ammonia.

Most of the more recent theories are founded on changes which the blood itself undergoes, as a direct consequence of the elevated temperature to which it is subjected at the time the injuries are sustained. An early view, recently re-advanced,<sup>4</sup> is, that owing to the loss of serum, the blood is so much thickened as to be unable to pass through the capillary

<sup>1</sup> Follin, *Traité de path. externe*.

<sup>2</sup> Falk, *Virchow's Arch.*, Bd. 53.

<sup>3</sup> Catiano, *Virchow's Arch.*, Bd. 87.

<sup>4</sup> Tappeiner, *Centralb. f. d. med. Wissensch.*, 1881.



vessels into the venous radicles. Overheating of the blood,<sup>1</sup> with associated paralysis of the heart, due to shock from irritation of a large area of nerves, is by some held accountable for immediate death, and reflex diminution of vascular tonus for death which, though not immediate, is not long delayed.

A number of observers attribute early death from burning and scalding to destruction of, or to interference with, the function of the red blood corpuscles by the heat, but they differ in their interpretation of the mode in which the injury to the corpuscles occasions death. Some hold that destruction of the blood corpuscles liberates fibrin ferment<sup>2</sup> which causes coagulation of the blood in the heart and other organs, and so gives rise to infarcts in the kidneys, liver, and bowels, causing subsequent erosions in the latter. Others<sup>3</sup> trace these changes to the debris of the broken down corpuscles, in support of which theory they instance the presence in the blood of numberless small coloured particles derived from the hæmoglobin, and of 'shadows' which represent the stroma of the red corpuscles, and, further, that the kidneys demonstrate the existence of free hæmoglobin by excreting it in the urine along with peculiarly coloured casts. Death is supposed to result from acute nephritis with uræmic poisoning. Part of the debris of the blood corpuscles is supposed to disappear in the pulp of the spleen and in the marrow of the bones. Against<sup>4</sup> this view it has been urged that the actual number of red corpuscles is not generally diminished after severe burns, but that loss of functional activity of a large percentage of them is thus occasioned, death resulting from the red corpuscles for the most part being rendered incapable of conveying oxygen to the tissues, the condition being that of acute functional oligocythæmia. In opposition to this is the statement that blood supposed to be thus disabled, on being shaken with air, takes up oxygen as easily and as copiously as normal blood.<sup>5</sup>

Still more recent views<sup>6</sup> corroborate and enlarge the theory of blocking of the blood vessels, especially of the small arteries, veins, and capillaries, which is not due to embolism, but to thrombosis, from clinging of the altered red corpuscles to each other and to the walls of the vessels, aided by the presence of a vast number of blood plates and debris of disorganised corpuscles. Salvioni,<sup>7</sup> whilst agreeing with this view, explains the formation of the thrombi in a different manner. He states, as the result of experimental observation, that the blood plates are so affected by the heat to which they have been subjected, that they deposit themselves on the walls of the vessels and give rise to the formation of minute thrombi, which are subsequently detached and are carried along by the blood, forming an enormous number of embolic nuclei. The ultimate pathological condition is due to three causes—the presence of the minute emboli, narrowing of the smaller vessels from excessive local heat, and, from the same cause, to an adhesive condition of the red corpuscles. According to Salvioni, the blood plates are not derived from other altered elements of the blood, but are normal and pre-existing. As proof of the important part played by the blood plates in the formation of thrombi, he states that by repeatedly defibrinating the blood in a living dog, it is rendered poor in blood plates, and that animals so treated survive the severest scalds, because no embolic nuclei are formed.

The preponderating opinion as to the cause of early death from burns and scalds is that it is a primary blood disorder, which chiefly results from injuries received by the red corpuscles when subjected to intense heat. Silbermann draws an analogy between the pathological changes which are caused by burning, and those produced by certain poisons—such as potassium chlorate, toluylenc-diamin, pyrogall, and anilin. Pfeiffer<sup>8</sup> attributes early death from burning to shock, or to the formation of a poisonous substance which resembles snake-poison in character, it is neither a ptomaine, nor yet a pyridin-base. This toxine is developed from the cell-material altered by the heat, it has nothing to do with the early blood-changes, and it belongs neither to the homolytic nor the agglutinating groups. The serum, or the sterile urine, from a rabbit

<sup>1</sup> Sonnenburg, *Verbrennungen*, *Deutsche Chir*, 1879

<sup>2</sup> Foà, *Rivista Sperimentale*, 1881

<sup>3</sup> Ponfick, *Berlin klin Wochenschr*, 1876 77, 1883

<sup>4</sup> Lesser, *Virchow's Arch*, Bd 79

<sup>5</sup> Hoppe Seyler, *Zeitschr f physiol Chemie*, Bd 5

<sup>6</sup> Welti, *Beitrage, zur path Anat*, Ziegler u Nauwerck, 1889, Silbermann, *Virchow's Arch*, 1890, Bd 119, Kleb's *Handbuch d path Anat*

*Arch per le scienze mediche*, 1891

<sup>8</sup> Virchow's *Arch*, 1905

which has survived burning is toxic to rabbits, mice, and guinea-pigs in amounts which are harmless when derived from a normal animal

It has frequently been observed that children succumb to burns and scalds more easily than adults. This is explainable on the ground that the child's skin is thinner, and, consequently, exposes the blood more freely to the action of the heat, and also that the resistance of the red corpuscles to any adverse influence is less in childhood than in adult life

When death takes place immediately in the presence of fire, especially within a building, it may result from poisoning by CO or CO<sub>2</sub> developed by combustion. If from CO, the blood and muscles may present the characteristic cherry-red appearance, and the blood may yield the absorption spectrum indicative of carboxy-hæmoglobin, if from CO<sub>2</sub>, the blood and muscles will be dark in colour

**Post-mortem Appearances—External.**—The appearances vary according to the severity of the injuries and the length of time the patient has survived them. In many cases the manner in which the injuries were caused produces characteristic indications. A burn caused by radiant heat leaves a white appearance of the skin. Contact with flame singes the hair, and blackens the skin from deposition of carbon, if not of great severity, there may be blisters at or round the site of the burn. The flame of an explosive, such as a mixture of coal-gas and air, scorches and mummifies the skin

Gunpowder-explosions not only blacken the surface from deposit of carbon, but, unless the patient is too far removed from the explosive, particles of unexploded powder are driven into the skin. The same observation applies to coal-mine explosions, after which the body is frequently blackened all over and profusely tattooed

Red-hot solids or molten metals produce effects which vary according to the length of time they remain in contact with the surface. If for a short time the skin only may be disorganised, in which case there will probably be blisters or the remains of blisters close to the burn. If the burn is very severe the soft structures will be roasted or carbonised like over-cooked meat, in such cases there will probably be an entire absence of blisters. From this degree up to combustion of the entire body—nothing but a few fragments of bones being left—all stages may be met with

The parts of the body protected by clothing are less injured by fire than those which are exposed. In slighter cases the covered parts escape altogether. The converse may occur if the clothing itself takes fire, which is notably the case when it is saturated with an inflammable liquid, such as petroleum, burns thus caused are of exceptional severity, and are further characterised by the distinctive odour of petroleum

**Scalds** caused by steam or boiling water produce appearances which are characterised by the presence of blisters—which, however, may be absent at parts owing to stripping of the cuticle—and by the hairs not being affected. Superheated steam produces a dirty-white, boiled appearance of the surface, which has lost all elasticity and feels sodden, in such cases blisters may not be seen. Scalds caused by liquids which have higher boiling points than water leave proportionally more severe indications

If the patient has lived some days, there will probably be signs of reaction which modify the original appearances

The bodies of those who are found dead from exposure to great heat are usually contorted, the limbs being in a state of flexion, so that, when the body is lying on the back, the arms project either upwards in a defensive attitude,

or across the chest. This condition is due to **heat-rigidity**, the causation of which was explained when the subject of cadaveric rigidity was discussed.

**Internal Appearances.** The **brain** is often shrunk to little more than one-fourth of its original volume, its form being fairly well maintained. When the head has been exposed to the direct action of flame, or other source of high temperature, in such a way as to char the scalp down to the bone, a localised extravasation of blood, which is frequently brick-red in colour, may occasionally be found on the corresponding inner surface of the skull, between the bone and the dura mater. This occurs with an intact cranial vault, and according to Strassmann<sup>1</sup> is caused by the heat driving the blood from the inner surface of the cranial walls. Littlejohn<sup>2</sup> records a case of this kind in which the dead body of a woman was found with the head on the bars of the fire-grate. The scalp was charred and the skull was exposed and blackened by the heat, but not injured otherwise. On removing the skull-cap a large brown blood-clot and a quantity of whitish-brown fatty material were lying on the dura mater, over the frontal and temporal areas corresponding with the position of the burnt spot. The **heart** is usually filled with blood. The **lungs** are shrunk, and often of a reddish colour. Particles of soot may be found in the larynx, trachea and bronchi, sometimes the mucous membrane is found injected and covered with froth, which may exist in amount sufficient to fill the air-passages. The **kidneys** may show signs of nephritis, and on section may present the appearance of reddish-brown markings, due to filling of the straight tubules with debris from the blood. This sign is not constant, but is sometimes very well marked. Frankel<sup>3</sup> records three cases of death from burning in which it was present, together with degenerative changes in the epithelium of the glomeruli and convoluted tubules. The mucous membrane of the **stomach** and **bowels** has been found to present a reddened appearance with swelling of the follicles. Curling<sup>4</sup> was the first to direct attention to the occurrence of ulcers of the duodenum in cases of death from burning, when the patient had survived some time, they have been frequently observed, although they are not present in the greater number of cases, Wilkes,<sup>5</sup> out of 37 cases, failed to observe any symptoms of the duodenum being affected, of these cases 12 were examined post-mortem. These ulcers are probably due to thrombosis of the small artery that supplies the part. They are more common in women than in men, whereas the idiopathic duodenal ulcer is much more frequent in men.

The **liver** shows nothing distinctive. The **uterus** may be found but little changed, although the corpse is almost consumed, the **testicles** also resist the action of fire. The **blood** frequently presents a peculiar and characteristic appearance, being of a cherry-red colour, very like that of blood from a case of carbon-monoxide poisoning. As previously stated, it is not uncommon for the blood of persons who have been burnt to death to contain carbon-monoxide, but the condition now under consideration, although the naked-eye appearances resemble those due to carboxy-hæmoglobin, has nothing to do with it, the absorption bands are those of oxyhæmoglobin, and reduction can be effected with the usual reagents. Falk<sup>6</sup> attributes the light colour to coagulation of some of the albumen of the blood in microscopic coagula, which diminish the absorption of light by reflecting a portion after it has traversed only a thin superficial layer of blood, the colour, therefore, appears brighter from a purely physical cause, no chemical change in the colouring matter taking place. In

<sup>1</sup> *Offiz. Bericht d. preuss. Medicinalb. Ver.*, 1898

<sup>2</sup> *Edin. Med. Journ.*, 1899

<sup>3</sup> *Deutsche med. Wochenschr.*, 1899

<sup>4</sup> *Med. Chir. Trans.*, 1842

<sup>5</sup> *Guy's Hospital Reports*, 1856

<sup>6</sup> *Vierteljahrsschr. f. ger. Med.*, 1888

fresh blood, this coagulation occurs at  $62^{\circ}$  to  $63^{\circ}$  C, in blood that is undergoing decomposition, it either does not occur at all, or else requires a higher temperature, due to complete or to partial conversion of the albumin into peptone, which does not coagulate with heat. Cherry-red blood from this cause is produced both in the bodies of those who have died from burning, and also in dead bodies which have been exposed to a sufficiently high temperature, on the other hand, cherry-red blood due to the presence of carboxy-hæmoglobin indicates that the individual in whose body it is found was alive whilst the fire was in progress. The urine of those who have died from burning frequently contains sugar—a positive reaction therefore would be corroborative of that mode of death if the victim was not previously glycosuric.

The most important question that the medical jurist has to answer after having examined a body injured by fire is—Did the burning take place **during life or after death?** The **external** signs, to which attention is directed with the view of answering this question, are limited to those displayed by the skin. When the injury is slight and is confined to the surface (a large area of the skin being affected), it presents the appearance during life of a diffuse **erythema** with patches where the epiderm has been detached. This appearance, produced equally by dry heat and by boiling water, is due to active hyperæmia, and can only occur during life, it is never found on a dead body that has been exposed to fire. Unfortunately it disappears, as a rule, after death, leaving the skin either pale or discoloured with post-mortem stains, it may sometimes be found, however, near parts that have suffered more severely. With certain reservations, the presence of **blisters** indicates that the injuries were received during life, to fulfil the indication the blister should contain fluid rich in albumen, and it should be surrounded by an injected margin either limited to its immediate circumference or blending into the diffuse erythema above mentioned. When the cuticle is detached the base of the blister, if examined soon after death, will be red, it is to be noted, however, that on the one hand, this redness of the base often disappears after death and on the other, that a slight reddening of the base of a blister produced after death may take place in consequence of exposure to air, it is, therefore, an indication of no positive value.

According to Hofmann, a blister having *all* the characteristics of those produced during life cannot be produced ten minutes after death. If an appropriate degree of heat is applied to the dead body a blister may be raised, but it will be filled either with aqueous vapour which condenses after withdrawal of the heat—the cuticle collapsing—or with fluid which contains little or no albumen, at the most only sufficient to cause opalescence on boiling it in a test-tube, whereas the fluid from a blister raised during life affords evidence of abundance of albumen on being similarly treated. Blisters are more easily produced on dead bodies that are œdematous, but, of course, the signs of vital reaction are wanting. Taylor and Stevenson record the case of a man who was drowned, whose body (with the object of resuscitation) was placed in a hot bath within a few minutes of the accident, no pulsation being perceptible, blisters containing bloody serum were produced by the high temperature of the water although the man was apparently dead when placed in the bath, and the attempts at resuscitation were unsuccessful. With regard to the occurrence of blisters on the bodies of those who have been exposed to the action of dry or moist heat, the inference to be drawn is—that if the fluid contained in the blisters is rich in albumen, and the blister is surrounded by an injected margin, it was either produced during life or *immediately* after death.

When the body has been subjected to a still higher temperature there will

be no blisters. In this case, the margins of the burns may afford some clue, if with the aid of a lens, or microscope, the capillaries of the corium can be distinguished as a blackish-brown network, the inference is that the burns were produced during life, burns produced in the dead body do not *usually* exhibit this network. The indication, however, is to be utilised with reserve, as a similar appearance may be produced by burns on those parts of the surface of the body where post-mortem staining is present, and again it has certainly been absent in undoubted cases of burning in the living. When the soft parts are completely charred it is impossible to say whether the burns were produced before or after death.

The **internal** differential signs of burns on the body produced before or after death are few in number, and not very distinctive in character. The blood may afford a clue if that obtained from the heart contains much CO, the presumption is that the individual from whose body it is derived was living when the injuries were sustained, because the hæmoglobin in bulk only combines with CO when passing through the lungs whilst they are respiring the gas, the conditions necessary for producing the combination are obviously wanting in the dead body. The naked-eye appearance of the blood must not be accepted as proof of the presence of carboxy-hæmoglobin, direct comparison of its spectrum with that of oxyhæmoglobin and evidence of non-reduction, or only partial reduction, after the addition of a reducing agent are required to substantiate its existence. Particles of soot or carbon in the larynx or the trachea are indicative of respiration having taken place whilst the fire was in progress.

Death by burning is usually **accidental**. This results either from isolated causation limited to the individual, as when the clothing catches fire, or from general causation, such as occurs when an inhabited building takes fire. From the nature of their clothing, women are more liable than men to fall victims to accidental burning of the first-named type, the fabrics of which dresses are frequently made are very inflammable, and a passing contact with fire is sufficient to kindle them into a blaze, when the victim with an instinctive impulse to flee from danger rushes about and thus fans the flames. Accidents of this kind have not unfrequently happened to gulls in ball-dresses, which usually are of exceptionally inflammable material, and before the flame is spent, or is extinguished, fatal mischief generally results. The upsetting or the explosion of paraffin lamps is another source of danger. Children are frequently burnt by playing with fire or with lucifer matches.

Considerable differences exist in the mortality of boys and girls from burning, as shown by the following figures compiled from the Registrar-General's Reports for 1906 to 1911, in England and Wales —

Ages	Boys	Girls	Ages	Boys	Girls
0 to 1,	214	234	4 to 5,	368	775
1 to 2,	443	466	5 to 10,	389	1,427
2 to 3,	840	623	10 to 15,	47	369
3 to 4,	580	729	15 to 20,	33	261

It will be seen that there is a marked drop in the mortality of boys after the age of three, whereas in girls the rate continues to rise until the age of five, the fall which then occurs being neither so rapid nor so great. This difference is undoubtedly due to the difference in the style of clothing worn by the two sexes. A series of petticoats, skirts, and pinafore separated by intervening layers of air, is a highly inflammable arrangement. Among boys the drop in the mortality occurs just at the age when male attire is substituted for this

costume. On the other hand, it is not until years of discretion are reached that a decline is shown among girls, and even then the mortality continues to be much higher right through the ages of adolescence.

In recent years the Registrar-General has endeavoured to tabulate separately deaths from burning where flannelette clothing was being worn, and deaths where the clothing consisted of other material. In a considerable number of cases, however, information as to the nature of the material was not reported. The following tables show the mortality at different ages in the two groups for the year 1919 -

#### FLANNFLETTE CLOTHING

Ages	Boys	Girls	Ages	Boys	Girls
0 to 1,	3		4 to 5,	2	14
1 to 2,	3	5	5 to 10,	2	23
2 to 3,	5	2	10 to 15,		3
3 to 4,	4	8	15 to 20,		4

#### OTHER CLOTHING AND NOT DESCRIBED

Ages	Boys	Girls	Ages	Boys	Girls
0 to 1,	3	8	4 to 5,	16	43
1 to 2,	13	8	5 to 10,	13	86
2 to 3,	31	21	10 to 15,	6	37
3 to 4,	18	30	15 to 20,	2	36

These tables clearly show that in death from burning in children, the style of the clothing is a much larger factor than the nature of the material of which the clothes consist.

Accidental scalding is not uncommon in children of a tender age, caused by the child pulling over a tea-kettle or pot containing boiling water, the contents of which are discharged over its body. Internal scalding is occasionally produced by children drinking directly from a tea-pot, the contained liquid being very hot.

Isolated cases of accidental burning and scalding in male adults occur chiefly in the pursuit of occupations attended with special risk—such as metal-founders and brewers.

Accidental fatal burning of general causation is more frequent in hotels, theatres, and other public buildings than in private houses. When theatres and public halls full of people take fire, a large percentage of deaths is due to suffocation from mechanical compression of the chest mutually produced by a number of persons seeking exit at the same door, and to the inhalation of the oxides of carbon.

When **homicide** has to be taken into consideration in respect to a dead body that is found partially consumed, the victim almost invariably has been killed by other means, fire is only made use of to conceal the crime. If the question, "Was death caused by the fire?" can be answered in the negative, suspicion of homicide is aroused, unless evidence to the contrary is forthcoming. When the body is not too far consumed, indications of violence may be visible—such as wounds produced by firearms, fractures of the skull, incised wounds, and marks of strangulation. It must be remembered, however, that stones, bricks, or slates not unfrequently fall from buildings which are on fire, and they may strike the head of a living or dead person and fracture the skull, the fire itself also may produce fissures or fractures of the cranial bones. Injuries closely resembling incised wounds may be produced by extreme heat, usually,

but not invariably, they occur at the flexures of the joints, in a case observed by Culling, the clefts in the skin and underlying structures were traversed by vessels and nerves in such a way as to preclude their causation by a cutting instrument. A strangulation-mark on more than one occasion has been clearly visible on a body that was almost carbonised. Schuppel<sup>1</sup> examined the carbonised remains of a boy ten years old, whose body was found in a burning cottage, and discovered indications of homicidal strangulation in the form of a distinct groove encircling the greater part of the neck, well-differentiated from the blistered and charred surface above and below it. The preservation of the mark is owing to the constricting medium being allowed to remain round the neck—it protects the skin underneath it, if the cord is removed before exposing the body to the fire, the mark will probably be destroyed.

The identification of bodies partially consumed by fire is easy or difficult in proportion to the degree of injury they have sustained, the lines on which such an investigation is to be conducted are laid down in the section on personal identity in the dead. The very considerable diminution in size and weight undergone by a body that has been partially carbonised must be borne in mind. The remains should be carefully searched for metallic objects which may afford evidence for identification, such as keys, rings, watches, and more especially, gold plates to which artificial teeth are attached.

### PRETERNATURAL COMBUSTIBILITY.

Under ordinary conditions the human body is with difficulty consumed by fire, a high temperature has to be maintained for a considerable time before the soft structures are carbonised. Exceptionally, an exactly opposite condition exists—the body is so easily consumed that the term “spontaneous combustion” has been applied to it. It may be premised at once that there is no evidence whatever to justify the use of the word “spontaneous,” but there can be no doubt that an extraordinary high degree of combustibility occurs in rare instances to which the term **preternatural combustibility** would more correctly apply. The peculiarity of the phenomenon consists in the fact that bodies which acquire this unnatural combustibility burn without the aid of the heat derived from the consumption of combustible matter other than afforded by the tissues themselves, the wooden floor on which such a body consumes is merely charred over the area that is in contact with it. This is totally opposed to the normal condition, in which the soft structures of the body not only refuse to burn of themselves—that is, by the simple application of a light—but they demand the consumption of an amount of fuel many times in excess of their weight before they can be destroyed, and when partly consumed soon cease to burn if the fire is withdrawn. It is to be inferred, therefore, that in preternatural combustibility some exceptional chemical change takes place, by which products of higher combustibility are developed than those which normally exist in the human body.

The subjects of this exceptional condition are usually fat, bloated, and much addicted to alcohol, exceptions have been met with in which the victims were spare and temperate of habit. They are almost invariably beyond middle age. The usual history is that the individual was in the midst of a debauch when the event took place, and that he was last seen some hours before, more or less profoundly under the influence of drink. The close relation between

<sup>1</sup> *Vierteiljahrschr f ger Med*, 1870

alcohol and preternatural combustibility of the body has led to the supposition that it might be the immediate cause, but this has been proved by experiment to be impossible. Chassagnol<sup>1</sup> steeped dead bodies in alcohol, and injected it into dogs' veins without increasing the combustibility of the tissue. Anyone who has thrown into the fire an old pathological specimen, which has been preserved in spirit, may have noticed that after the alcohol is burnt off, the solid tissues frizzle and burn slowly away, much as they would do in the recent state. Preternatural combustibility is evidently due to the formation in the body of some substance which is capable of burning alone when once ignited, not in a smouldering way, but with a luminous flame, as is proved by several cases in which flames were seen to proceed from bodies thus burning. It is exceedingly improbable that this substance can be any of the solid tissues, or fluids which have undergone chemical change, there is no analogy for such a complete alteration of property taking place within the living or the dead body.

It seems most likely either that exceptionally early and abnormal processes of decomposition of the tissues set in, or that, in some other way, inflammable gaseous products are formed, which, on escaping, become accidentally ignited, and the heat produced by their combustion consumes the tissues. A suggestive case is reported by Gull<sup>2</sup> of a large, fat, bloated drayman, of intemperate habits, who during cool weather was admitted into Guy's Hospital at nine o'clock one evening, and died an hour or two after admission. On the following day at one o'clock no signs of ordinary putrefaction were present, but the body was remarkably distended all over, the skin and the parts beneath being filled with gas, which, in the absence of any signs of decomposition, was suspected to have been exhaled from the blood, when punctures were made through the skin, the gas escaped and burnt with a flame like that of carburetted hydrogen, as many as a dozen flames were burning at the same time over the distended body. If this man had died in his own house, he would probably have passed his last hours alone, in a small room with a fire in it, and what so likely as that the gaseous emanations from his body should have become ignited, with the result that another case of "spontaneous combustion" would have been reported. A typical instance of preternatural combustibility is related by Archer<sup>3</sup>. An elderly woman of very intemperate habits, who was a large consumer of spirits of all kinds, lived alone in a small house. One morning, smoke was seen issuing from the closed shutters of her sitting-room, and, on breaking into the house, a small pyramidal heap of broken, calcined human bones, on the top of which was a skull, was found on the floor in front of a chair. All the bones were completely bleached and brittle, every particle of soft tissue had been consumed, and yet a table covered with a baize cloth, within three feet of the remains, was not even scorched. The rest of the furniture in the room was also intact. It is significant that the ceiling and the upper part of the walls of the room were scorched, evidently the result of flame.

In cases of so-called spontaneous combustion, there is nothing to show that the subject was alive at the time the burning commenced. On the contrary, the indications are rather against such a supposition, if alive, the victim must be in a profound state of alcoholic coma, or else there would be instances recorded of cries for help, which are wanting. The formation of inflammable gases in the body, moreover, is not incompatible with life. Beatson<sup>4</sup> records the case of a man who had foul eructations. He got out of bed one night and

<sup>1</sup> *Bull. Soc. de Chirurgie de Paris*, 1874.

<sup>2</sup> *Med. Times and Gaz.*, 1885.

<sup>3</sup> *Brit. Med. Journ.*, 1905.

<sup>4</sup> *Brit. Med. Journ.*, 1886.



struck a match in order to see the time, whilst blowing out the match his breath took fire, and exploded with a report sufficiently loud as to awaken his wife. This is by no means a solitary case. The formation of inflammable gases in the digestive tract has been demonstrated by M'Naught<sup>1</sup> to be due to bacillary fermentation, which also accounts for the abnormal formation of such gases in the dead body. Thus, in the condition known as "foaming liver," through the agency of a micro-organism—*bacillus aerogenes capsulatus*—the liver soon after death is found to be distended with gas, which escapes on section and ignites if a light be applied. Adams<sup>2</sup> records a case in which the condition of the abdomen led him to believe that the bacillus commenced growing in the tissues the day before the patient died. It has been noticed in all cases of preternatural combustibility, that the trunk is the original site of the combustion, the head and the extremities usually escaping, or, at the most, suffering from the contiguous heat. In a case reported, with a history of the subject, by Reynolds,<sup>3</sup> the abdominal wall was charred completely, and there was a large hole about eight inches long in the middle line, the face was crimson as from fire, but not blistered, the hands, stretched above the head, were unburnt, the arms were burnt and blistered, but not blackened, the thighs were burnt to the bone as far as the knees, where the burning abruptly ceased. It is evident that the combustion started in the abdomen, and in this instance spread farther down than is frequently the case. The woman was lying on her back with the thighs and knees well flexed, so that the former would be brought into close contact with the abdomen, and would, therefore, be subject to the full play of the flames, the effect of the heat on the anterior muscles of the thighs would be to shorten them, and thus, in the first instance, to draw the thighs more closely towards the abdomen. The burning had taken place either before or immediately after death, as blistering with signs of inflammation was present on the surrounding parts.

It seems probable, that in cases of preternatural combustibility, inflammable gases are formed in the abdomen, either during life or from abnormal changes which commence immediately after death, that the gas is accidentally ignited, and that its combustion raises the temperature of the soft tissues, especially the fat, so high that they become carbonised and give off gases of an inflammable nature which also take fire.

The part played by alcohol in the process has yet to be determined—whether its prolonged abuse leads to abnormal metabolic processes or to post-mortem changes in the tissues, more especially in those of the digestive tract, by which inflammable products are formed, or whether it undergoes some peculiar decomposition and itself furnishes the inflammable gas, is a matter of conjecture. The close alliance, shown by statistics, between the prolonged abuse of alcohol and an unnatural degree of combustibility of the body, makes it very probable that they stand in nearer relation to each other, as cause and effect, than that which has been allotted to them by some authorities—namely, that the alcohol simply stupefies the victim and makes him incapable of self-rescue.

The subject has an important medico-legal bearing. A murderer after killing his victim may endeavour to conceal his crime by burning, or partially burning, the body, in some instances the surroundings might lend colour to the supposition that the case was one of preternatural combustibility, and of absence of criminal intervention—an allegation which has been made under

<sup>1</sup> *Brit Med Journ.*, 1890

<sup>2</sup> *The Montreal Med Journ.*, 1896.

<sup>3</sup> *Med Chron.*, 1891.

these circumstances. The points to be remembered are that in cases of true "self" combustion, there is rarely much indication of the effects of fire beyond that displayed by the body itself, and that the trunk chiefly suffers, the extremities for the most part being preserved. The complete destruction of tissue occasioned by "self" combustion could not be accomplished, in the case of an ordinary cadaver, without the expenditure of a considerable amount of extraneous fuel, evidences of which would be forthcoming, and in this case the effects would scarcely be limited to or most apparent in, the trunk.

### BURNS PRODUCED BY CORROSIVE FLUIDS.

Burns produced by corrosive fluids which come under the observation of the medical jurist are the result of malicious throwing of the fluid on to the face or other parts of the body of the victim, the substances used are the mineral acids and strong solutions of the caustic alkalies, of these sulphuric acid holds the first rank, so much so that a special name is given to the act "vitriol throwing". None of these substances will directly occasion death when thrown upon the body, unless under very exceptional circumstances, they may produce great disfigurement, however, and may indirectly conduce to a fatal issue.

As regards efficacy, it is not without reason that sulphuric acid is the substance usually made use of, it attacks organic matter energetically, and its corrosive action is only partially arrested by wiping the surface on to which it has fallen which is usually the only counteractive treatment available for the moment. **Sulphuric acid** in a concentrated form produces a **brownish-blackish** eschar if thrown on to the surface of the body, it is not necessary, however, for the acid to be of full commercial strength to effect this, but if it is much diluted, the mark produced will be grey. **Nitric acid** leaves a **yellow stain** or slough, the colour being caused by the action of the acid on albuminoid bodies, forming xanthoproteic acid, the stains of nitric acid sometimes appear brownish on the dead body. **Hydrochloric acid** may or may not leave a **whitish-grey** stain, it is not so destructive to tissue as are sulphuric and nitric acids.

There are certain appearances common to the eschars produced by all three acids. The surface is smooth and soft, in contradistinction to the irregular and hard surface of a heat-burn, and the surrounding zone produced by congestion of the small vessels of the cutis, which encircles the eschar produced by fire, is wanting in that caused by acids.

The **caustic alkalies** have a solvent action on albuminoids and fats. Potash-lye is especially corrosive in its action on the tissues, under its influence they become tumid, and the skin communicates a greasy feeling to the touch, subsequently the part becomes dry and resistant, the colour being dark, especially on the dead body. As is the case with the acids, the eschars produced by alkalies have no encircling zone of reaction.

If the surface is examined immediately after the receipt of the injury, direct evidence may possibly be obtained as to the nature of the corrosive, by brushing the surface with a small camel-hair brush (quill-mounted) dipped in distilled water, and then pressing the fluid out of the brush into a test-tube. The proceeding may be repeated once or twice, and the resulting fluid tested, first as to reaction, and then, if acid, as to the kind of acid, the appearance of the mark on the skin and the stains on the clothing will probably give a clue as to which acid should be sought for in the first instance. The same remarks apply to

injuries produced by alkalies, except, of course, as regards reaction and subsequent testing. This plan, however, is rarely successful, as the part in all probability will have been well laved with water before an opportunity is afforded of trying it. The character of the stains severally produced on fabrics, with the method of testing, will be found described in the section on Toxicology.

In the absence of direct proof obtained from the injured part, it is of the highest importance to ascertain the nature of the substance thrown from evidence yielded by the clothing, as the *attempt* to injure by throwing a corrosive, even should no bodily injury be sustained, constitutes felony. In the absence of bodily injury, the evidence required from the medical witness is an answer to the question— Was the substance thrown of a corrosive nature?

## CHAPTER XXIII

### MECHANICAL INJURIES AND WOUNDS.

ALMOST all injuries to the body produced by mechanical violence are comprehended, in the legal sense, under the title of "wounds." The definition is enlarged beyond that given by the surgeon by the words which follow in the statute "*Whosoever shall, by any means whatsoever, wound or cause any grievous bodily harm to a person*." The last clause of this sentence obviates the necessity for defining whether a given injury is or is not a wound, which was a question that formerly led to much discussion at almost every trial relating to matters of personal violence, and which not unfrequently facilitated the escape of a guilty person.

If the injury is not obviously of a very severe kind, medical evidence will probably be required to determine whether the life of the injured person is in danger or not, and whether "*grievous bodily harm*" has been inflicted. These questions can only be answered after a thorough examination of the injuries has been made, the appearances found being interpreted with the aid of an appeal to general experience. The subject involves the consideration of injuries, both in their general aspect, and also as to special characteristics dependent upon the kind of instrument with which they were produced.

### CONTUSIONS.

This is a comprehensive term, embracing all degrees of injury produced by blows or sudden pressure with objects that do not divide the skin, from a simple bruise that merely causes rupture of a few blood-vessels in the corium, up to complete disorganisation of the soft structures underlying the seat of injury. One result common to all contusions is the extravasation of blood. The amount effused is not exclusively determined by the severity of the blow—the anatomical structure of the parts injured exercises a considerable influence, tissues which are open in texture allow the blood from the ruptured vessels to distribute itself more freely than those which are firm, a blow in the region of the eye causes wider spread extravasation than one of equal force on the palm of the hand. The condition of the blood-vessels is another factor, when diseased, as in purpura, subcutaneous hæmorrhages occur spontaneously or on the slightest

pressure. In the absence of disease, many women develop bruises with the least possible pressure, this is a point to be remembered when estimating the degree of violence that has been exercised in the case of women of flabby constitution with soft skins, in whom a comparatively gentle grasp of the arm will produce bruises indicating the points of pressure caused by the thumb and fingers that might easily be attributed to excessive violence. When blood is effused in or immediately beneath the skin the discoloration produced is blue-black in colour. Pepper<sup>1</sup> directs attention to a change in colour not unfrequently undergone after death by superficial subcutaneous extravasations, which at first are bluish-black and subsequently become pink or scarlet.

**Superficial ecchymoses** make their appearance within a few minutes after their causation. **Deep-seated ecchymoses** may not appear for days, and then not always directly over the seat of the injury. If blood is effused from one or two relatively large vessels at some distance from the surface there may be no discoloration of the skin, but only a sense of fluctuation on palpation.

In certain parts of the body injuries of the severest kind may be inflicted without producing external signs of bruising. The wheel of a cart may pass over a man's abdomen and rupture the liver and other internal organs without the least trace of ecchymosis being visible in the skin, either before or after death. A kick or blow may rupture the bladder or intestines without leaving any external mark.

Superficial ecchymoses that are undergoing absorption display a surrounding zone of colours—brown, green, and yellow, fading into the normal skin-tint, are usually present. These appearances, due to modifications undergone by the hæmoglobin, are initiated at the periphery, and gradually advance towards the centre of the discoloured spot, affording evidence of wider distribution of the effused blood than was at first apparent. At some distance from the ecchymosis, the skin that was not discoloured when the bruise was first produced takes a lemon tint, which is continuous with the deeper shades of the original discoloration. Ecchymoses under the conjunctiva being very superficial, and being backed up by the white sclerotic, are of a bright red colour, and when fading pass through no chromatic changes except to yellow. **Deep-seated extravasations** of blood show no superficial colour changes other than a gradual diminution in intensity.

## WOUNDS.

Under this head are comprised incised wounds, punctured wounds, and lacerated wounds.

**Incised wounds** are those made with cutting instruments or with objects presenting more or less sharp edges, such as pieces of broken glass or crockery-ware. The length is usually greater than the depth. A characteristic of an incised wound made during life is that its edges retract and curve outwards, causing the wound to gape. This is to a great extent due to the elasticity of the skin, aided or counteracted, within limits, by the direction of the underlying fibres of the connective tissue and muscles, in accordance as to whether they run parallel with, or transversely to, the line taken by the incision, in the limbs the course of the fibres runs lengthwise, except near the joints. The cleanness and evenness of the edges of an incised wound are determined by the manner in which it was produced, and to a still greater extent by the nature of the object that produced it—a sharp knife will divide the skin in a clean, regular

<sup>1</sup> *The Lancet*, 1887.

way very different from the cut made by a sharp-edged stone. When a sharp knife is resolutely used for the purpose of inflicting a bodily injury, the conditions most favourable for the production of a clean incision are fulfilled—a suitable instrument appropriately handled. By falling on a sharp-edged stone, it is quite possible for a man to cut himself very severely, but the conditions are much less favourable as regards the regularity of the wound especially of its margins. The skin being very elastic tends to recede before the edge of a cutting instrument, necessitating a sweeping or sawing movement in order to divide it, any such application of the edge of a stone or broken piece of earthenware would be of fortuitous occurrence. When incised wounds are really caused by falling on broken pottery, or the like, the broken edge is driven through the skin with but little, if any, of the movement necessary to divide it cleanly, the consequence being that the margins of the wound have a more or less contused appearance. To enable this to be distinguished the wound must be examined shortly after its infliction before any inflammatory changes take place. The distinction between an incised wound made with a dull-edged knife with notched blade and one made with a sharp stone may not be so easy.

Some emphasis has been laid on the appearance of wounds made by sharp cutting instruments as compared with those caused by the wounded person falling on broken pottery and the like, because in cases of alleged criminal wounding with cutting instruments the defence usually advanced is that the prisoner with his fist struck the prosecutor, who fell on a sharp stone which inflicted the wound. It is true that in either case the prisoner would be guilty of wounding, but the use of a knife is held greatly to increase the culpability of the act.

In certain parts of the body where the skin is somewhat stretched, with little between it and the bone underneath—as is the case over the malar bone, for example—the violent tangential impact of a rounded hard body, such as a cricket-ball or even the clenched fist, may cause a wound with difficulty distinguishable from one made by a cutting instrument. A blow on the scalp with a poker or policeman's truncheon may produce a similar effect. Such wounds are due to sudden, forcible stretching of the integument, splitting or tearing it open. Due reserve should be displayed in attributing wounds in this and other similar parts of the body to the use of a cutting instrument, the possibility of the above described causation should ever be borne in mind. When such a wound is seen immediately after it is made, minute examination with a lens may reveal irregularities of the margins inconsistent with its production by a cutting instrument, while in the deeper parts of the wound little bridges of tissue or vessels may be found.

The amount of **hæmorrhage** from incised wounds depends on the number and size of the vessels divided, it is usually considerable. In a recent wound blood-clots will be found, and the connective tissue laid bare will be infiltrated with blood.

**Punctured Wounds.**—These are wounds made by a piercing instrument and have usually a depth greater than their length. The risk to life caused by punctured wounds is determined by their situation, the depth to which they penetrate, and the size and shape of the weapon by which they are produced. A fine exploring needle may be repeatedly introduced into certain organs—such as the liver—with impunity, whereas a still finer instrument introduced elsewhere may produce fatal results. Magnan<sup>1</sup> relates the case of a woman who committed suicide by passing an ordinary pin, barely an inch and a quarter

<sup>1</sup> *Comptes Rendus de la Société de Biologie*, 1890

long, into the left side below the breast at the sixth intercostal space, on examination after death seven small punctures were found in the apex of the heart, and the pericardium contained between seven and eight ounces of blood.

A punctured wound made with a sharp-edged and pointed knife leaves a wound through the skin having almost parallel but slightly concave margins which meet at acute angles at the two extremities. This is invariably the case if the knife or dagger is double-edged, and usually so with an ordinary blade—that is, sharp on one edge only—unless it is of considerable thickness at the back. When the blade is thick at the back throughout its entire length, or when not so thick, if plunged in up to the hilt where the back is thickest, the opening is wedge-shaped—one extremity forms the apex and the other the base of an exceedingly acute triangle. With either a double- or single-edged weapon, if it is thrust in and withdrawn in the same plane, without lateral movement, the aperture will be slightly less in length than the breadth of the instrument that produced it, this must be borne in mind when comparing the size of a punctured wound with a suspected weapon. For the same reason it is risky to express a decided opinion as to the depth of a stab wound in the living subject from a comparison between the length of the wound through the skin and the width of the blade at a given distance from the point. The discrepancy in size between the wound and the instrument that produced it is due to the elasticity of the skin which may also cause the edges of the skin, in a wound produced by the thrust of a knife or dagger, to be everted—the skin gripping the blade as it is withdrawn. If the weapon is not thrust in and withdrawn in the same plane the wound through the skin will be longer than the breadth of the blade, as any movement of an oscillating nature divides more skin than is necessary for its direct passage. If a part of the body is transfixcd by a long-bladed weapon, the secondary orifice will be smaller than the primary, and its edges will be everted.

Punctured wounds made with pointed weapons of circular transverse section—such as a butcher's steel—are not circular in outline. Such instruments act as conical wedges, they do not cut but they split the tissues, the consequence being that instead of a round aperture one resembling a slit is produced, the direction of the long axis of which is determined by the course of the fibres which enter into the structure of the tissues perforated. It is to be noted that the course of the fibres in the superficial structure of an organ does not take the same direction over its entire surface, in some organs the fibres radiate, in others they are arranged somewhat circularly from one or more centres. Punctured wounds caused by sharp-pointed fragments of glass and similar objects are characterised by a combination of cutting and splitting of the skin and the tissues perforated. The margins of the wounds will be jagged, and will bear traces of bruising, the appearance being different from that of a wound produced with a sharp-pointed knife. Punctured wounds of a similar type are sometimes caused by certain artisan's tools, the pointed end of a file that is intended to be driven into a wooden handle has blunt cutting edges, which, when the implement is used as a dagger, or is accidentally thrust through the skin, tear open rather than cut the tissues. Sometimes the wound made by an instrument which has a special form is sufficiently distinctive as to betray the kind of weapon used.

The danger to life from punctured wounds is frequently determined by the loss of blood which results, the external hæmorrhage may or may not be profuse, but when only a little blood flows out of the wound, severe internal hæmorrhage may be taking place, and may speedily cause death.

**Lacerated wounds** are those which are produced by objects that tear the tissues in place of cutting them. Such a wound may be sustained by an individual who falls violently on a hard projection— as the corner of a box or step, or by catching against a nail or pointed hook with a sudden movement that rips open the surfaces. Lacerated wounds may be produced by blows with an obtuse-angled weapon, as a hammer or poker, by the teeth of man or of the lower animals, by crushing, and a variety of other ways.

The characteristics of lacerated wounds are their irregularity and the jagged, swollen appearance of the margins. When produced by violent impact with a blunt-edged object, the skin in the neighbourhood of the wound is ecchymosed, the underlying tissues are more or less disorganised, and if a bone is near the surface, it is likely to be fractured. Heavy falls on a flat surface, such as the flags, a macadamised road, frozen ground, or the ice, may cause lacerated wounds— a causation frequently alleged to account for those suspected to be due to criminal violence. Some parts of the body are more likely than others to be lacerated in this manner, the scalp over the eyebrows, the tissues over the malar bone, or those over a flexed joint, as the knee, for example. It is often difficult or impossible to say whether such wounds were directly caused by a blow or indirectly by the fall which resulted from the blow. The site of the wound may enable a decision to be arrived at, if it is at the vertex it is more likely to have been caused by a blow than by a fall, unless the wounded person fell from a height, the presence of soil or grit in the wound lends probability to the supposition that it resulted from a fall. Not only is the wound to be carefully examined, but the clothing also, especially the hat if the head is the part injured. When the head-covering shows a tear corresponding to the wound, and the margins of the tear are coated with gut or other substance derived from the ground, the wound being similarly coated strong evidence is afforded that the injury was the result of a fall.

If a weapon by which the wound is alleged to have been made is forthcoming, it should be examined for marks of blood, for hairs, and for fibres derived from the clothing of the injured person. A minute comparison must be made of the weapon with the wound in order to determine whether such a weapon could have caused the wound. Sometimes the implement used leaves special indications in or about the wound—as, for instance, soot off a poker, and rust off a hammer or an iron bar.

Lacerated wounds do not, as a rule, bleed so freely as incised wounds, but if the structures lacerated are very vascular, bleeding to a fatal extent may occur. This is especially the case with regard to the external female genital organs, death from hæmorrhage has resulted from a lacerated wound of the vulva caused by a violent kick from a foot armed with an iron-bound clog, or even with a thickly-soled boot.

**The distinction between wounds inflicted before and after death.**—The characteristics of **bruises** produced during life have already been described. Marks of a similar kind may be produced immediately after death, and, further, local violence inflicted in the last moments of life may not produce any obvious change in the colour of the skin until after death has occurred, therefore, it must not be assumed that the presence of a bruise-mark proves survival for a time after the casual violence was inflicted. There is no practical difference between a bruise produced immediately before death and one produced immediately after, as effusion of blood from ruptured vessels of the skin may occur for a short time after death—that is, whilst the blood is fluid and warm. Absence of cardiac pulsation prevents the occurrence of extravasation of blood

to any great extent after death, but in subcutaneous ecchymoses the difference in amount is too insignificant to enable a decided opinion to be expressed as to their ante- or post-mortem origin. The appearance of a bruise inflicted during life cannot, however, be produced on a dead body after the lapse of a short interval after death. If an individual has survived the infliction of injuries for a day or two, the usual colour changes at the margins of bruises will be visible.

**Incised wounds** made after death differ from those inflicted during life, in respect to absence of gaping and of much hæmorrhage, a short survival adds another indication—tumefaction of the edges of the wound, still longer survival leads to changes, due to reparative processes, which remove all doubt as to whether the wound was produced before death. Gaping being due to contractility of the skin, areolar tissue, and muscles, is not an absolute sign of infliction during life, the edges of an incised wound made *immediately* after death will retract and cause the wound to gape, the extent of retraction is less than during life, but, since it occurs at all a certain reserve must be maintained in the expression of an opinion. Within a very short time after death the contractility of the tissues is lost, an incised wound produced at or after this period presents the appearance of a slit, the edges of which lie closely approximated without any trace of eversion. On separating the edges, little or no blood will be found effused, unless a large vein has been divided, that which is present may be either fluid or more or less coagulated, any infiltration into the areolar tissue is slight as compared with that which takes place in a wound inflicted during life. If, in the course of an incised wound inflicted during life, an artery of moderate size is included amongst the structures divided, a considerable amount of hæmorrhage takes place, if the wound is made after death, the bleeding will be relatively insignificant, being limited to the veins, therefore the amount of blood found on the body and clothing, and on the surface on which the body lies, may afford a valuable indication. Spurting of blood to a distance from a divided artery is a sign that the wound was inflicted during life, but it only occurs with small arteries, with large arteries the blood-pressure is lost at a short distance from the cut end, and the blood is not projected far. The appearance presented by blood spurted from a small artery on to a wall or piece of furniture is that of a succession of small spots forming a line, or possibly a confused mass with isolated spots around it, if a larger artery spouts on to an object close at hand, there may be a splash of blood which, as long as it remains sufficiently fluid, trickles down the surface and forms a vertical and more or less linear mark.

**Punctured wounds** made before or after death present appearances respectively analogous to incised wounds, a punctured wound made after death may (from perforation of a large vein) cause a considerable quantity of blood to be poured out into one of the cavities of the body.

**Lacerated wounds** made *immediately* after death may present appearances like those seen in wounds inflicted during life, if produced a short time after, the possibly ecchymosed margins will not be swollen.

When making a **post-mortem examination** of a body on which there is a **wound**, the first thing to do is minutely to examine the part without disturbing anything, and to write down at the time all the points observed. The external appearance of the wound, its size and position, the amount of blood poured out, and if the body is seen where it was originally found, the presence of blood-stains from spurting on neighbouring objects, all claim special attention. If there is blood observe if it is coagulated, and if so, whether firmly or not. The



condition as to cadaveric rigidity and post-mortem stains is always to be investigated in medico-legal cases. The interior of the wound is to be examined, first, as to the presence of clots, blood, and staining of the connective tissue, and then its direction must be ascertained. In the case of a stab wound a blunt bougie may be cautiously passed in, but the operator must be prepared to answer a question that he may be subsequently asked as to whether by doing so he did not extend the wound. With due care there is no risk. The deeper part of the wound is to be reached by dissection in the direction taken by the bougie, if possible without interfering with the external opening. This is especially important if the weapon that is supposed to have caused the wound is not forthcoming, it may subsequently be found, in which case a comparison of it with the outer wound is very desirable. The depth of the wound and the structures through which it passes are to be noted, the possible presence of foreign substances not being overlooked. If a bone is injured, it may be well to remove it (or the injured portion of it), as corroborative evidence.

## CHAPTER XXIV

### SPECIAL WOUNDS AND INJURIES.

#### INJURIES OF THE HEAD AND SPINE.

THE scalp is one of the regions where wounds resembling those caused by a sharp cutting instrument may be produced by blows from rounded or obtusely-angled objects, in this way a flap may be detached leaving a portion of the skull bare. In the absence of unhealthy inflammation uncomplicated scalp wounds are not dangerous to life, any difficulty in prognosis is usually caused by the possible presence of complications in the form of injury to the skull or the brain.

Injuries to the **skull** are dangerous to life either remotely or immediately the former is the case when a blow on the head is followed by inflammation of the diploe, producing septicæmia two or three weeks after the receipt of the injury, in the meantime the patient appears to be doing well. Immediate results follow **extravasation** of blood within the cranium, either between it and the dura, in the arachnoid or the pia mater, or in the brain substance itself, the symptoms are those of cerebral compression—slow, laboured breathing and pulse, total insensibility with absence of reaction of the pupils, which may be either dilated, contracted, or unequal. When such an extravasation is the direct result of a blow on the head the skull is usually fractured. Occasionally a case occurs in which a man who is not suffering from marked degenerative changes of the vessels is knocked down and, without the skull being fractured, shortly after develops the fatal symptoms just enumerated, in such cases medical opinion is required as to a possible causal relation between the blow and death. The transverse sinus most frequently suffers, more rarely hæmorrhage from traumatic violence without fracture takes place in the pia and arachnoid. This mode of death not unfrequently happens to habitual drunkards and to persons advanced in life, in both, the vessels are liable to spontaneous rupture, which may occur at the time the injury was received. In one case

of this kind the deceased was in a paroxysm of passion at the time, and was about to resort to extreme violence when his adversary struck him a blow which caused him to stagger and fall, afterwards, insensibility came on, which ended in death, on examination cerebral hæmorrhage from a degenerated vessel was found, but no signs of injury to the skull. In such cases allowance must be made for the condition of the deceased at the time, and without expressing a positive opinion, it is to be admitted that in elderly people and in habitual drunkards cerebral hæmorrhage may be determined by excessive mental excitement. Post-mortem signs of injury to the head, though without fracture, would point to local violence as the cause of the hæmorrhage. It is obvious that a mixed causation exists in many cases, a vessel is predisposed to rupture, and does rupture from a degree of violence that would not injure one in a normal condition. In such cases the medical witness must explain the condition of the parts and content himself with so doing, unless he has good reason for thinking that the rupture did or did not result from the blow.

The question is sometimes further complicated by apparent recovery from the effects of the blow or fall, followed, after an hour or more, by relapse into unconsciousness which terminates in death. The defence naturally is, that death was due to an independent cause, the interval of recovery showing that the blow could not have caused rupture of a vessel, else there would have been no return to consciousness and power of walking. This inference, however, is contrary to fact. In many instances cerebral hæmorrhage from both traumatic and idiopathic causes has commenced, producing a certain degree of unconsciousness, the bleeding has temporarily ceased, the brain has recovered from the pressure, and the patient has regained consciousness, renewed bleeding from the same source has then ended fatally within a few hours.

**Concussion of the brain** is another way in which external violence may cause death without the skull being fractured. As generally understood, concussion of the brain does not cause disorganisation, therefore, no indications as to the cause of death are revealed at the necropsy. In fatal cases the injured individual is suddenly rendered unconscious, respiration is irregular and fitful for a short time, and then ceases altogether, the pulse is probably imperceptible from the first, the surface is cold and there is entire absence of reaction. Less severe cases usually end in recovery.

**Contusion of the brain** is either circumscribed or diffused. In the former variety one or more centres of disorganisation, or of extravasation of blood occur, in the latter, the extravasations are distributed throughout the substance of the brain, and, possibly, also on its surface. The mischief may be limited to the site of the blow, or it may be situated at an entirely different part, owing to the so-called *contre coup*, the base and the middle lobes most frequently suffer on account of the irregular conformation of the base of the skull. The symptoms are those of cerebral irritation, there is usually unconsciousness with great restlessness and, possibly, tonic or clonic spasms. In a slighter form the primary symptoms may pass away and the patient may recover, but there is risk of inflammatory changes occurring, which may spread to the meninges and ultimately cause death. In other instances the patient recovers with partial loss of memory or with some degree of paralysis.

**Fractures of the skull** may be divided into those produced by blows with weapons presenting a small striking-surface, which may be either pointed, rounded, or flat, and those caused by violent contact with surfaces of larger area. Fractures caused by blows with a heavy weapon of **limited striking-surface**, such as a narrow-headed hammer, very frequently display the shape

of that part of the weapon which came in contact with the bone. If the striking-surface has a rectangular outline, and it strikes the bone in the vertical line with considerable momentum, the resulting fracture will also have a more or less rectangular shape, the portion of the bone on which the striking-surface falls being driven forward by the violence of the impact, the detached portion almost entirely absorbs the momentum with which the implement is endowed, so that comparatively little tension is put on the surrounding bone, which consequently escapes injury more or less completely. If the same implement reaches the skull with less momentum it may cause an irregularly depressed fracture with fissures traversing the surrounding bone. If the fracture, viewed from the outside of the skull, has a sharply-defined characteristic outline, the inner table will be found irregularly splintered off all round the aperture, which is consequently much larger on its inner than on its outer aspect. Fractures equally characteristic of the casual weapon may be produced by sharp blows with the spherical knob of a poker or the head of a life-preserver, the resulting injury takes the form of a circular concave depression when viewed from the

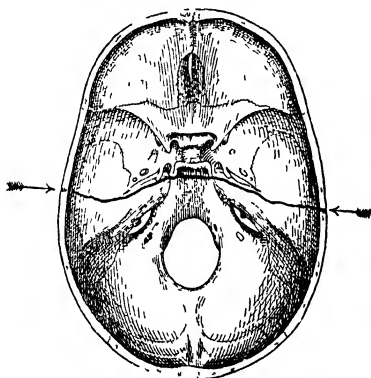


Fig 19 — Fracture from bi lateral compression

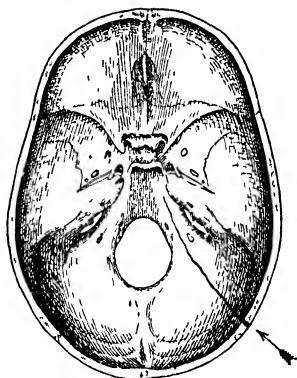


Fig 20 — Fracture from one sided compression

outside, and appears as an irregularly formed fissured projection on the inner surface. Penetrating injuries of the skull, made with sharp-pointed instruments having thin blades, like a dagger or a knife, take the form of the blade or else that of a depressed fracture, such as may result from a blow with any weapon having a very limited striking-surface.

The direction of fractures of the **base** and of the **vault** of the skull caused by blows or by sudden pressure with objects of **large contact-area** is determined by certain physical laws. Wahl<sup>1</sup> and Messerer<sup>2</sup> were the first to demonstrate the part played by the elasticity of the skull in the production of fractures of this kind, and their deductions, which are of the highest importance to the medical jurist, have been corroborated by numerous clinical observations.

<sup>1</sup> *Volkman's Klin. Vorträge* (Chirurgie No. 73), 1883

<sup>2</sup> *Exper. Untersuch. über Schädelbrüche*, 1884

Fractures of this kind are divided by Korber<sup>1</sup> into two groups —(a) Those produced by bi-lateral compression of the skull, and (b) those which result from violence applied to one side only, the head as a whole being free to recede from the impact of the blow. In both groups the line of fracture runs parallel with the axis of compression.

(a) Fractures in the first group are produced not only by interposition of the skull between two opposing forces, both of which are in motion, but also if the skull is simply prevented from receding when the blow is delivered, as happens when a blow falls from above on the head of a prostrate person, the head resting on the ground, or when a person who is standing with his back against a wall receives a blow on the forehead. In these cases the skull is compressed between two points or poles, the compression being greatest at the equatorial line, where the fracture begins and tends to spread towards each pole, in other words, it begins at the base or the vertex (more frequently the former) and spreads outwards in the direction of the two points where compression is applied, the fissure being widest at its starting point. Korber records, with drawings, a number of cases of fracture of the skull variously produced. In one instance, a woman was killed by being struck with a wooden mallet near the left ear while asleep with the head on a pillow, the line of fracture which resulted ran across the base from ear to ear. In another case a man lying asleep on the right side received a blow from a hatchet near the left eye, after death the skull was found to be fractured diagonally through the base, from the left orbit to the right parietal protuberance.

Among this group are those fractures which result from blows on the vertex (the individual being in the erect posture), the counter pressure being derived from the resistance offered by the vertebral column.

(b) When the compression is one sided, as when a man in the upright posture is struck with a blunt instrument on the side of the head, the head being free to recede under the momentum imparted by the blow, the fracture begins at or near the point of impact and travels in a direction parallel with the axis of compression, it rarely goes beyond the middle line, and tends to narrow in width the further it advances. For example, a man was struck on the left temporal region with a large stone and dropped down dead, a comminuted fracture was found where the stone struck the head, and a compression fracture running transversely across the left middle fossa, narrowing as it reached the sella Turcica, where it terminated.

A fracture of the skull in which a portion of the bone is driven in by a blow from a weapon having a limited striking-area may be accompanied by a compression-fracture, the impulse being sufficient to compress the skull as well as to force a fragment of it inwards. The compression-fracture may be situated at some distance from the spot where the blow fell, but it will run in a direction parallel with that of the blow.

In some instances the bones of the cranium are exceptionally thin, being no thicker at parts than stout paper, a slight blow is then sufficient to cause a fracture. The thickness of the skull should always be noted when examining the bodies of those who have sustained fracture of its component bones, excessive tenuity being frequently pleaded in criminal cases.

If a man standing in the erect posture is knocked down by a blow on the head, and dies in consequence, the skull being found to be fractured, the question may be asked. Is it not possible for the skull to be fractured by a fall on level ground? It is possible, provided that the ground is hard and unyielding.

Severe fracture of the skull with depression or other injury to the brain is not invariably followed by insensibility, and many cases have occurred where great damage has been done to the skull without depriving the sufferer of power of movement. Agnew<sup>2</sup> relates two striking examples. A man was run over by a tramway-car, he got up immediately after the accident, walked a short distance to his house, opened the door with a latch-key, went upstairs to his bedroom on the second storey and got into bed, where on the following morning he was found insensible with portions of the parietal and temporal bones deeply driven into the brain. In the second case, a man had half of the frontal bone

<sup>1</sup> *Deutsche Zeitschr. f. Chirurgie*, 1889.

<sup>2</sup> *Medical News*, 1887.

with a considerable portion of the pre-frontal lobes carried away by the bursting of a flywheel, he was dazed for a few moments only, and eventually made a good recovery. Such instances show the necessity for expressing a guarded opinion as to the loss of consciousness and power of locomotion in cases where a dead body is found with severe injury to the brain.

**Injuries of the Spine.**—The regions most liable to injury are the upper cervical, the lower cervical, and the upper lumbar. The extent of the injury sustained varies from a slight sprain up to fracture and separation of the bodies of one or more vertebrae. Slight sprains are often exceedingly difficult of detection, the symptoms being purely subjective, if the cord suffers, symptoms either immediate or remote supervene. **Hæmorrhage** into the meninges—*hæmatorrhachis*—usually produces sudden loss of power, violent pain at the seat of the hæmorrhage, which will probably radiate along the nerves given off from the affected segment, with clonic and tonic spasms, and subsequently more or less paralysis. The bleeding takes place either outside the membranes, within the arachnoid or, more rarely, beneath it. **Concussion** of the spine may be followed by delayed development of symptoms, several weeks elapsing before anything manifests itself more definite than the sensation of having undergone a severe shake. Then various paræsthesiæ develop, with difficulty of walking and especially of stooping, urination may be enfeebled from loss of expulsive power, and there may be partial paralysis of the legs. Along with these symptoms there are usually disturbances of the special senses, especially of sight and hearing, the mental condition deteriorates—evinced by loss of memory, inability to concentrate the ideas, and general irritability. These symptoms not unfrequently follow the shake produced by a railway collision, and it is no easy matter to determine how far they result from diffuse injury sustained by the cord and the effects of shock to the nervous system generally, and how far they may be simply of a subjective nature, due to the influence of “suggestion.” An accurate mapping out of the anæsthetic and hyperæsthetic areas with subsequent comparison with the known origin of the sensory nerves of the parts will probably show whether the sensory disturbances are really due to changes in the cord or are only of psychical derivation. The subject is too wide to be discussed in a text-book on forensic medicine, special works must be consulted. Much valuable information will be found in Morley’s *Injuries of the Back in their Medico-Legal Aspect*<sup>1</sup>

**Fractures** of the spine, with or without displacement, may be caused by direct or by indirect violence. The former results from the injured part being struck by, or striking against, some hard substance, as, for example, a blow with a bar of iron or a fall from a scaffold on to a projecting piece of timber. Fracture from indirect violence may result from forcible bending of the body forwards or backwards, it usually occurs in the cervical or dorsal regions. Forcible bending of the spine may be caused by external violence, as when a sack of flour falls from a height on the head of a man who is slightly stooping forwards and doubles him up, or by a sudden powerful voluntary muscular contraction, such as was made by an insane female patient in an asylum, who, to avoid the spoon by which she was being fed, violently jerked her head back and thus fractured the cervical spine. Moore<sup>2</sup> examined the body of a prisoner, aged about thirty-five years, who had delusions, in the middle of the night “he dived off the bed” on to the smooth tiled floor of his cell, the bed being only seventeen inches high. When seen, immediately after, the head was in a slightly flexed position, and the man was quite conscious. On the second day

<sup>1</sup> *Internal Congr. of Med.*, 1913.

<sup>2</sup> *Communicated*, 1901.

he died, and at the autopsy the fifth cervical vertebra was found to be broken into three or four pieces, the arch into two pieces, and the arch of the sixth was detached from its body, the cord being crushed. The vertebræ presented no appearance of disease, nor were they ankylosed.

The duration of life after fracture of the spine may be roughly stated to be in direct proportion to the distance between the injured part of the cord and the medulla, but much depends on the amount of displacement of the bones, if the cord is not compressed and had not been crushed, there is less immediate danger to life than when the converse is the case. A slight laceration of one of the intervertebral substances with displacement, so as to cause pressure on the cord without fracture of a vertebra, is of more moment than a simple fracture without displacement. The cord is more likely to be damaged by an injury to the spine resulting from indirect than from direct violence. Fractures in the lumbar or dorso-lumbar regions, although accompanied by displacement, are not necessarily fatal, recoveries have taken place after fracture with displacement, even in the cervical region, though, as a rule, such injuries are speedily fatal. From the medico-legal standpoint it is of some importance to remember that fracture with dislocation of the upper cervical vertebræ is not necessarily *immediately* fatal. It is usually assumed that, on account of implication of the phrenics, injury to the cord at or above the level of the third cervical vertebra is suddenly fatal. Eve<sup>1</sup> records a case of fracture of the odontoid process with forward displacement of the atlas which compressed the cord, and yet the patient lived two hours and a half. Gurling<sup>2</sup> had a case in which death did not ensue for twenty-eight hours after fracture of the first three cervical vertebræ, although the cord was injured at the level of the third.

**Injuries of the Face.**—Incised wounds of the face bleed freely, but, except when an artery such as the maxillary or lingual is divided, the hæmorrhage is not dangerous in a healthy individual. Blows with blunt weapons may fracture the nasal and malar bones, and cause "grievous bodily harm," so far as permanent disfigurement is concerned, but they are not dangerous to life, the possible occurrence of erysipelatous inflammation, however, is to be taken into account in giving an opinion as to the risk to life. Blows on the face may cause hæmorrhage into, or detachment of, the retina, in some cases atrophy of the optic nerve has followed a blow in the region of the eye, in both these conditions permanent blindness of the affected eye may result. The eyeball may be directly injured by the penetration of foreign bodies, and opacity of the lens or its capsule produced.

**Injuries of the Ear.**—Hæmatoma auris or, as it is also called, the insane or asylum ear, consists of an effusion of blood into the auricle. In the insane two factors are probably concerned in its causation—local hyperæmia due to degeneration of the vaso-motor fibres of the cervical sympathetic, and mechanical violence. A third factor vascular degeneration—may occasionally lead to the formation of hæmatoma auris after a relatively slight degree of violence, or in exceptional cases to its spontaneous formation. Robertson<sup>3</sup> states that the hæmorrhage usually takes place from new vessels in the walls of cysts which are the result of chronic degenerative changes in the ear cartilage. It is frequently met with in insane people, especially in asylums, and is probably sometimes caused by blows on the ear delivered by impatient attendants, the left ear generally suffers. In the early stage the auricle is swollen and tense,

<sup>1</sup> *St Barth Hosp. Rep.*, 1887

<sup>2</sup> *Lond Hosp Rep.*, vol 1

<sup>3</sup> *Edin Hosp Reps.*, 1896

with the appearance of a collection of blood under the skin, this may subsequently become organised, and distort and shrivel the ear. *Hæmatoma auris* is not limited to the insane, it may result from various acts of local violence, such as occur in football scrimmages, in wrestling and pugilistic matches.

The tympanum may be ruptured in a variety of ways, of which one only, as occasionally giving rise to legal proceedings, need be particularised, it is when the injury is alleged to have been caused by a box on the ear. As a rule, it is the left tympanum that is ruptured, there will be no external bleeding, unless other injuries have been inflicted.

## INJURIES OF THE NECK AND CHEST.

**Injuries of the Neck.** From the medico-legal standpoint injuries in this region may almost be paraphrased into "**cut-throat**," the greater number being of suicidal origin. When the close relation and superficial position of the numerous structures in the throat upon the integrity of which life depends is considered, and further, the fact that most of the wounds in this region are self-inflicted with the object of causing death, it is strange how often that object is defeated by the more immediately vital structures escaping injury. A large proportion of suicidal wounds of the throat stop short just before the important vessels are reached, the structures actually divided being of secondary importance as regards the maintenance of life, it is owing to this fact that so many cases of cut-throat are seen in the wards of hospitals, if the large vessels are divided, death is almost certain to take place before surgical aid can be obtained.

Wounds in the neck are usually incised wounds or stabs, the former being most common. On account of the looseness of the skin, especially in old people, it is not unusual for more than one incision to result from a single stroke of the knife. Incised wounds of the throat vary to the fullest possible extent, from mere nicks through the skin to gashes that well-nigh sever the head from the trunk. Punctured wounds of the neck are very dangerous from the risk of perforation of one or more of the large vessels. At the back of the neck a thin-bladed knife may be passed between two of the vertebræ and with very little external mark may produce death from injury to the cord.

**Injuries of the Chest.**—Sharp severe blows on the chest near the cardiac region, inflicted by objects of large area, may cause sudden death without leaving any trace of injury, blows with instruments of less dimensions are likely to cause fracture of one or more of the ribs. A common way of accidentally fracturing the ribs is by a fall against a projecting object, as the corner of a table, compression of the chest between two opposing forces is also a frequent cause. Very exceptionally, ribs are fractured by coughing, sneezing, and by sudden movements of the arms in endeavouring to preserve the equilibrium of the body when the feet slip whilst walking. **Stabs** of the chest, which do not penetrate the thoracic cavity, as a rule are not dangerous, although there may be considerable bleeding from the mammary and thoracic arteries. Penetrating wounds are dangerous, but not necessarily fatal, unless a vital part has been injured. If the **lungs** are injured, emphysema, pneumo- or hæmo-thorax may ensue, as well as secondary results—pleurisy or pneumonia. The danger lies in the liability to profuse hæmorrhage, which may be speedily fatal, although severe penetrating wounds of the chest, followed by copious hæmorrhage, may be survived. A case came under the care of Hulke<sup>1</sup> in which the patient fell

<sup>1</sup> *The Lancet*, 1888

off some "high steps" whilst she had a picture covered with glass in her hands, the glass was shattered by the fall, and when the woman was lifted up a large fragment of it was found sticking in her back, the withdrawal of which was followed by much bleeding. The wound on the back, at the level of the tenth rib, and three inches to the right of the spine, was three inches long, the tenth rib was divided. The fragment of glass had transfixed the thorax, for in front there was another wound three-quarters of an inch long opposite the posterior wound, but higher, being in the seventh intercostal space. The woman recovered.

Laceration of the lungs, followed by hæmorrhage, may be produced by external violence without fracture of the ribs. A boy was run over by a cab, and died the following day, the ribs were uninjured, but the lung was extensively lacerated, and the pleural cavity was full of blood.

Penetrating wounds of the **heart** are usually fatal, but not necessarily immediately so, when death is not immediate, the duration of life varies from an hour up to many months, and, in exceptional cases, even years. West<sup>1</sup> records the case of a man who was stabbed with a knife, profuse hæmorrhage and great collapse following, from which he recovered and survived four years, after death, a linear cicatrix was found in the wall of the right ventricle, probably the cavity of the ventricle was not reached. The ventricles, especially the right, are more frequently wounded than the auricles, the left auricle being rarely penetrated. It might be supposed that immediately fatal hæmorrhage would more frequently follow penetrating wounds of the auricles than of the ventricles, on account of the thinner walls of the former. Statistics,<sup>2</sup> however, do not support this supposition. Coats<sup>3</sup> reports the case of a girl, aged ten years, who fell on to an iron railing, one of the spikes of which penetrated the right side of the chest where the fourth rib joins its cartilage, the patient survived nine days. Examination after death revealed an elongated wound in the pericardium, three-quarters of an inch long and one-quarter of an inch broad, opposite to which was a wound through the wall of the right auricle. Strassmann<sup>4</sup> relates the case of a man who was stabbed with a knife in the left mammary region, the wound traversed the fourth intercostal space, through the left ventricular wall—where it was a quarter of an inch wide—into the cavity, the patient lived four days, and on the day following the infliction of the injury he lifted heavy weights. At the autopsy about three and a half ounces of blood were found in the pericardium and about ten ounces in the thoracic cavity. Slight superficial wounds of the heart may prove speedily fatal. In a case recorded by Thompson,<sup>5</sup> a man died within a few hours from the insertion of an ordinary pin, one inch and a half long, through the space between the fifth and sixth ribs in the area of the cardiac impulse, the anterior wall of the left ventricle was wounded, but its cavity was not penetrated, the pericardium contained seventeen and a half ounces of blood.

It is important to remember that those who have sustained wounds of the heart may perform certain actions, although the injuries are speedily fatal, the following are instances selected from a number of cases of wounds of the heart collected by Fischer.<sup>6</sup> One individual, after being thus wounded, ran 450 paces, another mounted several steps, a third walked a mile and a half, and a fourth (in whom the right ventricle, coronary artery, lung, diaphragm,

<sup>1</sup> *St Thomas's Hosp Rep*, vol 1, N S

<sup>2</sup> See Tables in Holmes & Hulke's *System of Surgery*, third edition, vol 1

<sup>3</sup> *Glasgow Med Journ*, 1891

<sup>4</sup> *Trans of the Royal Acad Med Ireland*, 1888

<sup>5</sup> *Lehrb der ger Medicin*, 1895

<sup>6</sup> *Langenbeck's Archiv*, ix



liver, stomach, spleen, and colon were wounded with a sabre) made ten steps before he fell

In exceptional instances myocarditis, or rupture of the left heart, may be caused by a blow on the chest from a blunt instrument, without either the ribs or the skin over them being damaged. Heidenhain<sup>1</sup> reports the case of a sailor who was thus injured by a heavy blow on the chest from a capstan bar. Such injuries when followed by myocarditis may soften the cardiac musculature, and lead to its rupture many days or weeks after the receipt of the injury, probably no external signs of injury would be visible after death. Immediate rupture of the heart may be caused by violent blows, as the kick of a horse, or by the passage of a cart-wheel over the chest, without any external signs of violence. In such cases the right side of the heart, most frequently the right auricle, is the seat of the rupture. In relation to immediate rupture of the heart or of the commencement of the aorta after a blow delivered on the chest, it is to be remembered that atheromatous ulceration, or the presence of a diminutive aneurismal dilatation, may predispose to spontaneous rupture. A man<sup>2</sup> was admitted under the author's care in the Salford Royal Hospital in 1892, who, whilst stooping, suddenly felt faint, and subsequently was profoundly collapsed, he rallied, and three days after, whilst in bed, he was again attacked in the same way, he rallied once more, but six days after the second attack he was found dead in bed. At the necropsy the pericardium was seen to contain a large amount of fresh blood, and there were signs of recent pericarditis with much organised fibrin. A small rupture through the base of an ulcer had taken place at the commencement of the aorta, microscopical examination showed a gradual thinning of the coats down to the aperture, probably due to necrosis ulceration from thrombosis. The weak spot gave way at the moment the patient was first attacked temporary occlusion, with subsequent renewal of the hæmorrhage, occurred on two occasions, the last being speedily fatal. The patient was a vigorous, healthy looking man, and had followed a laborious employment up to the moment he was first attacked, had the rupture taken place after a push or slight blow of the fist, it might have been misinterpreted as proof of extreme violence. Rouse<sup>3</sup> records a somewhat similar case of a man, aged fifty two, who, whilst playing football, suddenly fell down in a sort of faint, he recovered and continued to play, but after the game was over he again became faint and collapsed. He revived and lived for a week, when he had another attack and died in less than five minutes. At the necropsy a rupture, half an inch in diameter, was found in the left ventricle, the pericardium containing twelve ounces of blood, fatty degeneration of the heart and of most of the other organs was present.

## INJURIES OF THE ABDOMEN.

Death may suddenly result from a blow delivered by a blunt weapon on the pit of the stomach, in such cases it is probable that reflex paralysis of the heart is the cause of death. Maschka<sup>4</sup> records two such cases- in one, a boy was struck with the fist over the stomach, in the other, a strong man was struck over the same region with the flat part of a shovel, both died at once, and in each case the result of the necropsy was negative. Beach<sup>5</sup> records the case of an intoxicated man who was arrested in the street by the police, he resisted so violently that one of the officers struck him a blow with his "club" in the epigastric region, when he suddenly became quiet and powerless, and on arrival at their destination he was found to be dead, not the least trace of injury could be discovered, neither externally nor internally.

Rupture of the abdominal viscera may result from violence without any external sign of injury. The stomach is rarely ruptured, if it is, the pyloric end and greater curvature generally suffer. The intestines are not unfrequently ruptured by external violence, according to Weil, the commencement of the jejunum and then the ileum most frequently suffer. Such accidents, probably, are invariably fatal, although the dangerous symptoms may not develop for some time, and meanwhile the injured person may walk a considerable distance

<sup>1</sup> *Deutsche Zeitung f. Chirurgie*, 1895

<sup>2</sup> *Trans. Path. Soc. Manchester*, 1892

<sup>3</sup> *The Lancet*, 1892

<sup>4</sup> *Vierteljahrsschr. f. ger. Med.*, xxxi,  
Wiener allg. med. Zeitung, 1884

<sup>5</sup> *Medical News*, 1882

Even when the blow is inflicted with a formidable object such as a horse's hoof, there may be no external marks. Hunter<sup>1</sup> saw a boy who had been fatally kicked in the abdomen by a horse, there were no ecchymoses nor signs of injury externally, but the jejunum was bruised and ruptured. F. J. Smith<sup>2</sup> relates the instructive case of a girl aged seven who was accidentally knocked down by a drunken man. She had immediate abdominal pain which suggested rupture of the intestine, but when the abdomen was opened no rupture was found. Two days afterwards, death occurred, and at the autopsy acute peritonitis was found, without trace of tear, bruise, or wound of the intestine, nor of any of the other viscera.

The **liver** is frequently ruptured by heavy pressure, such as a wheel transmits from a loaded vehicle, the rotating movement of the wheel favours the absence of signs of external injury, and the liver may thus undergo extensive rupture without the skin of the abdomen being ruffled or discoloured, rupture from a blow or kick also may occur without external bruising. Death usually results, and it may do speedily, from hæmorrhage into the abdomen, but in spite of copious internal hæmorrhage, the patient sometimes lives several days, after slight ruptures recovery may take place. A ruptured liver does not necessarily prevent movements being made by the patient after receipt of the injury.

Rupture of the **diaphragm** is not common apart from other and severe injuries to the abdominal viscera. In itself it is not likely to be followed by immediate death, but will probably lead to serious results from protrusion of the stomach or bowels into the thoracic cavity. Laceration of the diaphragm usually occurs in consequence of falls from a height or from compression of the trunk by a cart-wheel passing over it. It is generally met with on the left side.

Rupture of the **spleen** is usually fatal. What has been said about the liver applies equally to the spleen. In persons who have lived in tropical countries and who have had intermittent fever the consequently enlarged spleen has been ruptured by a very slight push or blow. Brouardel<sup>3</sup> states that, during a scuffle, a man was slightly pushed against a bureau, he felt nothing at the moment, but shortly after he became pallid and died, at the autopsy the spleen was found ruptured. Pellereaux<sup>4</sup> gives the history of thirteen cases of rupture of the enlarged spleen, of these five were spontaneous, the others being due to apparently inadequate causes, such as a simple fall in the street or out of a carriage. External indications of local violence are absent in such cases.

Extensive rupture of the **kidneys** is always fatal. Slighter rupture may be recovered from. Much bruising of the organs without rupture is likely to be followed by suppuration, which may cause death at a more or less remote period from the receipt of the injury.

Rupture of the **bladder** is a subject of special importance to the medical jurist, because, unlike some of the previously described injuries, it may be caused by a blow of the fist or a kick that would be inadequate to reach less accessible organs. It is an accident likely to occur to a drunken man, who, on account of the amount of liquor drunk and its benumbing effect on the nerve-centres, often goes about with a full bladder. In this condition a fall downstairs may be sufficient to cause rupture. The injury is almost invariably fatal when the intra-peritoneal portion of the bladder is ruptured, and but few recoveries take place when the extravasation of urine is outside the peritoneum. Bartels<sup>5</sup> states that 93 out of 94 cases of intra-peritoneal rupture died, and 26

<sup>1</sup> *The Lancet*, 1885

<sup>2</sup> *Lond Hosp P M Reports*, 1905.

<sup>3</sup> *Annales d'hygiène*, 1894

<sup>4</sup> *Ibid*, 1882

<sup>5</sup> *Langenbeck's Archiv* Bd 22

out of 63 in which the mischief was extra-peritoneal. Extensive rupture of the bladder is not necessarily prohibitive of locomotion, the severity of the injury being in some instances concealed for a time. Bartsch<sup>1</sup> relates a case in which a drunken man fell through a window twelve feet, on to hard ground. He was found lying on his side, and began to joke, making no complaints of being hurt, he was placed in an outhouse, where he lay for an hour, when he got up, went into the house and walked upstairs to bed. He rose the following morning, after having slept well, and took a walk of nearly a mile, he then had a rigor and returned to bed, after which he gradually became worse, and died sixty-three hours after the fall. On section no external injury was seen, but the peritoneum contained about 52 ounces of urine, and there was a tear in the superior and posterior wall of the bladder,  $2\frac{1}{2}$  inches in length, which went cleanly through the peritoneal coat. Eight hours elapsed before indications of the severity of the lesion manifested themselves. If, in the meantime, this man had quarrelled with anyone, and had in consequence received a blow, his antagonist might easily have been made responsible for his death.

A defence likely to be made in cases of rupture of the bladder from alleged criminal violence is that the organ ruptured spontaneously.

**Spontaneous rupture** of the bladder is of exceptional occurrence, even in those cases where a diseased condition of the viscus or of the urethra exists as a causal factor, which is capable of being recognised as such after death, the conditions alluded to are ulceration of the bladder-wall and organic stricture of the urethra. Paralysis of the bladder might also lead to spontaneous rupture, in such a case the previous history of the patient would help to explain its occurrence. Much is usually made by the defence as to the absence of signs of external injury, which, it is urged, proves that the organ ruptured from natural causes. This is fallacious—a full bladder—and the organ is always in this condition when ruptured by a blow which does not fracture the pelvis—is, of all the abdominal organs, the most likely to be ruptured from external violence without here being any outward signs of it.

Exceptional as is spontaneous rupture induced by previous pathological changes, a still rarer event has been recorded—spontaneous rupture without any explainable cause. Facewen<sup>2</sup> relates the case of a young man, aged nineteen, habituated to excessive indulgence in alcohol, who on one occasion, whilst nearly insensible from this cause, was taken to a common lodging house and put to bed. He remained in bed all the next day, on the evening of the following day he was drowsy and stupid and complained of pains in the abdomen, he died on the third day. After death the abdomen, free from external marks of violence, was found to contain a large quantity of straw coloured fluid, of which the bladder also contained a small amount. At the junction of the upper and middle thirds posteriorly an aperture existed in the bladder which would admit the tip of the little finger. There were no indications of disease or of ulceration or gangrene, neither was there any peritonitis, the urethra was healthy and free from stricture or obstruction—a No. 10 sound found its way into the bladder by its own weight. Brown<sup>3</sup> records another case of spontaneous rupture of the bladder without the presence of a stricture, the urethra allowing

No. 10 catheter to pass. After death the posterior part of the bladder was found coated with soft lymph, and about an inch below the reflection of the peritoneum, slightly to the right, was an aperture which admitted the forefinger with difficulty, the margins were rough, from the presence of recent inflammatory lymph, there was no evidence of ulceration or of pre-existing disease. Such cases are of extreme rarity, only one or two being recorded. It is very difficult to understand the mode of their occurrence, the two alternative conditions usually assumed are paralysis of the bladder of functional origin and persistent spasmodic stricture of the urethra.

**Incised and lacerated** wounds of the abdomen are not in themselves dangerous when the peritoneum and viscera escape injury, when forming a prognosis

<sup>1</sup> *Vierteljahrsschr. f. ger. Med.*, 1889.

<sup>2</sup> *The Lancet*, 1873.

<sup>3</sup> *The Lancet*, 1886.

the possible occurrence of peritonitis or of suppuration among the muscles and fasciæ must be taken into account **Punctured** wounds are necessarily dangerous if they penetrate the abdominal cavity, both from the chance of protrusion of the viscera through the wound, and, to a still greater extent, from the possibility of the viscera themselves being wounded **Small punctures** of the **intestines** are not necessarily fatal, the danger being proportional to the risk of their contents escaping to the peritoneal cavity, wounds of the small intestines are more dangerous than those affecting the large intestines Incised or punctured wounds of the **solid viscera** are dangerous in proportion to the hæmorrhage they occasion

**Injuries of the External Genitals.**—Injuries to the **male** organs are not unusually dangerous to life, although the bleeding is sometimes considerable Rupture of the urethra in the perineum leads to extravasation of urine into the connective tissue of the pelvis, which may be fatal unless surgical aid is promptly afforded

**Injuries to the Female Organs.**—**Incised** or lacerated wounds of the vulva and vagina often give rise to dangerous hæmorrhage, apart from parturition, they usually result from accident Birkett<sup>1</sup> relates two such cases A lady going into a dark room to micturate sat down on a water-ewer, the handle of which was broken off, leaving a sharp and jagged portion projecting about an inch, this produced a lacerated wound of the vagina, in the course of which the internal pudic artery was divided, death from hæmorrhage took place in about an hour after the infliction of the injury In another case, a woman was knocked down by a man, and died in a short time from hæmorrhage, the blood came from the vagina, in the wall of which was a wound extending towards the internal pudic artery At first it was thought that the woman had been stabbed with a knife, but it turned out that she had fallen on and broken a spittoon, a sharp-pointed fragment of which caused the injury In cases of criminal assault foreign bodies, such as pieces of stone or wood, are not unfrequently passed into the vagina, causing serious bruises and rents of the vaginal walls Violent sexual intercourse with young girls sometimes and with adult women may rupture the vagina, and occasion profuse and even fatal bleeding

The **unimpregnated uterus** is not frequently injured unless it shares in a general injury inflicted on the pelvic organs, the **gravid** organ, from its prominent position, is much more frequently wounded Penetrating wounds of the gravid uterus, whether caused by stabs through the abdominal walls, or by punctures inflicted by passing pointed instruments up the vagina in the attempt to procure abortion, are dangerous from the possible supervention of hæmorrhage, peritonitis, and septicæmia Kicks delivered against the abdomen of a woman advanced in pregnancy may produce miscarriage, or, by separation of the placenta, may give rise to serious hæmorrhage, violence of this kind may be followed by localised suppuration in the uterine wall, or by the formation of clots in the sinuses, which may cause death from embolism of the pulmonary artery The walls of the gravid uterus may be ruptured by blows, by pressure of the abdomen against hard projections, and by falls, and the fœtus may escape through the rent into the abdominal cavity Such accidents are usually fatal, though in some cases operative interference, and in others natural processes, have relieved the woman of the displaced fœtus, her life being spared Rupture of the uterus or vagina may take place during parturition, either in the course of natural labour, or during the performance of version or of

<sup>1</sup> Holmes and Hulke, *A System of Surgery*

extraction with the forceps. The medico-legal questions that arise relate to the reasonable amount of skill brought to bear in the treatment of the case. On the one hand, was artificial delivery performed with a reasonable amount of care and skill? and on the other—the uterus rupturing in the course of natural labour—ought artificial assistance to have been resorted to? Such questions are usually referred to obstetrical experts.

### FRACTURES OF BONES.

Simple fractures are not dangerous to life except in certain regions, as the skull and spinal column, from the medico-legal standpoint, the question is rather one of “grievous bodily injury.” The defence usually urged in criminal cases is either that the fracture resulted from accident, or, if acknowledged to have directly caused, that the injured bone was predisposed to fracture in consequence of disease or of some inherent abnormal condition.

Apart from diseases affecting the bones, which, from their nature, are sufficiently obvious, such as cancerous and sarcomatous growths, rickets, and mollities ossium, there are other general diseases and conditions which tend to render the bones more fragile than ordinary. It is well known that trophic changes occur in the course of certain diseases of the nervous system, amongst the structures liable to participate in these changes are the bones. Chemical analysis shows that the proportion of organic to inorganic matter is inverted as compared with healthy bone, the inorganic matter from diminution of the phosphates is reduced to less than half, and the organic matter in consequence of a large excess of fat is doubled. When these alterations in the composition of the bones are far advanced, spontaneous fracture of the long bones is not uncommon, when not so far advanced, a much less violent blow will fracture a limb than would do so in health. The diseases of the central nervous system, with which these trophic changes in the bones are chiefly associated, are locomotor ataxy and some forms of mental disease, especially general paralysis.

In some cases trophic changes in the bones appear to take place without the presence of any recognisable disease. Greenwood<sup>1</sup> states that a policeman taking part in a contest of throwing a cricket ball at some sports felt his arm snap in the act of throwing, on examination complete fracture of the humerus was found to have taken place at the lower third. The bones become brittle in old age from excess of inorganic constituents, and in young children they are liable to “greenstick” fracture from the converse condition. Ribs are occasionally fractured by coughing or sneezing. A curious case of apparently delayed completion of a fracture of the rib is related by Skyrme<sup>2</sup>. A man in getting out of an omnibus swung round and struck his right side against a projection, he felt some pain and tenderness, but went about as usual. Six days after, whilst sneezing, he felt something snap in his side, which was followed by severe pain, greatly increased by deep respiration, the tenth rib was found to be fractured at the junction of the bony and cartilaginous portions.

The medical witness may be called upon to state his opinion as to whether certain injuries to bones were the result of direct violence or not. A drunken man after quarrelling with another man is found to have sustained fracture of several ribs, and dies from pneumonia, his adversary is accused of having caused the injuries by blows, but denies that he struck the deceased, stating that he simply pushed him away, and that in his drunken condition he fell helplessly to the ground. Such a question is to be decided on general grounds,

<sup>1</sup> *Brit Med Journ*, 1890

<sup>2</sup> *Ibid*, 1891

the amount and character of external bruising, the position of the fracture, the number of ribs broken, whether fracture has occurred on both sides of the chest, together with other indications of violence more than can be accounted for by the simple act of pushing a man away. A similar question arises from time to time in regard to asylums and prisons, an attendant or a warder is accused of having caused the death of an inmate or prisoner by violence. The defence usually is that the injuries (often fracture of the ribs) were caused by a fall, or in the case of a prisoner recently taken into custody, that they existed at the time of his arrest. With regard to the latter point, it is possible for a man to go about with fractured rib or ribs without knowing that he has received any such injury. In the case of the insane the possibility of trophic changes in the bones must be borne in mind.

The following remarkable case is quoted from the Annual Report of the Board of Control for the year 1919 ---

"W B, male 42, a general paralytic. A few days after removal of left upper molar tooth by the medical officer a fracture of the lower jaw was diagnosed. Some six days later the patient died from pneumonia, and at the post mortem examination the following extraordinary conditions were found ---

"Fracture of lower jaw in front of the masseter on either side, old standing healed fractures of the 5th, 6th, 7th, 8th, 9th, and 10th, and a double fracture of the 11th ribs on the right side with strong fibrous union. Old standing fractures of the 7th rib (one fracture), 8th (four fractures), 9th (five fractures), 10th (three fractures), and 11th (three fractures), on the left side with fibrous union.

"The fracture of the jaw might, it was thought, have been caused by a sudden reflex contraction of the masseter muscles at the time of the extraction of the tooth, and the injury to the ribs could be stated, with probably absolute certainty, to have all existed for a considerably longer period than the patient's residence in the asylum. There was no history of accident or injury to the patient whilst he was in the asylum."

The length of time the fracture has existed may be of importance as regards determining the question of criminal violence or of accident, if it is clear that the fracture existed before the alleged violence was inflicted, the fact is, of course, greatly in favour of the accused. The processes which occur in the repair of fractured bones are well known, but their time-values are not capable of exact estimation. Apart from physiological variations, the age and the state of health of the injured person materially influence the rate of repair, the degree of injury sustained by the fractured ends, and the closeness or otherwise of their apposition have also to be taken into consideration. Within the first week after the occurrence of the fracture there will probably be nothing to be seen on post-mortem examination beyond effused blood, with more or less tearing or bruising of the contiguous soft structures, shortly after, indications of repair begin to manifest themselves. In simple fractures in which the ends of the bone have been kept in apposition from the first, but little, if any, 'provisional callus' is thrown out, it is mostly found in fractures of the ribs or of the clavicle—that is, in bones in which absolute rest cannot be attained. After fourteen or sixteen days, the blood at first extravasated will have disappeared or nearly so, the periosteum at the fractured ends will be very vascular, and beneath it and between the ends of the bone will be a number of cells proliferated from the osteoblasts amongst which calcification will be in progress. Under the most favourable conditions complete ossification does not take place, as a rule, in less than two months.

Evidence of **previous fracture** is usually not difficult to obtain when the examination is made **after death**. If on external examination the bone does not yield sufficiently clear indications, a longitudinal section of it will clear up

all doubts. In the **living** an old fracture may easily escape detection. If immediately after receipt of the injury perfect approximation of the broken ends of the bone was secured and maintained until complete ossification took place, especially if the part is deeply surrounded with soft structures, it will be quite impossible to recognise a remote fracture, if the fracture is recent it will be easier of recognition. In exposed situations, such as the front of the tibia, the difficulty is lessened.

The distinction between fractures produced in the **living** and in the **dead** subject is well marked, unless the fracture is produced immediately after death. If it takes place six or eight hours after death there will be no blood effused round the ends of the bone, unless a large vein has been divided, and, even then, the appearance will be distinct from that caused by extravasation from a number of vessels during life. A fracture caused immediately after death within a few minutes—may present an appearance indistinguishable from one inflicted during life.

### WOUNDS PRODUCED BY FIREARMS.

Wounds produced by firearms differ in appearance in accordance with the size and kind of projectile, the velocity with which it is endowed at the moment it impinges, the distance of the firearm from the body, and the angle at which it is presented.

Large bullets, other conditions being equal, produce more extensive wounds than those of less size. The old-fashioned spherical bullet is more apt to be deflected in its course than the modern cylindrical bullet with conical front, the wedge-like properties and higher velocity of the latter missile enable it to make its way through obstacles which would turn a spherical bullet to one side. A conoidal bullet endowed with high velocity causes infinitely greater damage than a spherical bullet, not only because it can penetrate a mass of tissues from side to side that would arrest a spherical bullet half-way, but also because its crushing action on the soft tissues and its power of splintering the bones are vastly greater.

When a small conoidal bullet endowed with high velocity passes through some of the soft structures of the body without striking any bone, the difference presented by entrance- and exit-wounds is not great, under ordinary circumstances, however, the difference is considerable. A large bullet traversing the body with less velocity produces a much more extensive aperture of exit than of entrance. If the weapon is fired point-blank the **entrance-wound** will be about the size of the bullet, and will have a more or less circular outline with torn edges which may form angular flaps, the margins will be ecchymosed and slightly inverted, except in parts where there is much subcutaneous fat, when they will probably be everted. If death is immediate, the surrounding skin may be pallid. The **exit-wound** will be larger and less regular in outline, with everted edges showing the subcutaneous fat. If the weapon is presented at an **oblique angle** to the body the entrance-wound will not present a circular outline, it will be more elliptical, the skin being ploughed up to one side. A bullet projected from a distance, if not too far spent to penetrate, will cause a larger and more lacerated entrance-wound than if it arrived with greater velocity. Only a limited value can be assigned to this as an indication of the distance from the body at which the weapon was fired, much depends on the kind of weapon, and on the amount and the energy of the explosive used.

When a firearm is discharged into the body at a distance of a few inches

only, the entrance-wound, in addition to the appearances already described, will be blackened and, possibly, scorched. The wound is diffusely blackened from the smoke of the explosion, and it is also tattooed by undeflagrated grains of powder being driven into it. The flame from the mouth of the firearm may singe the adjacent hair and scorch the skin. It may also set fire to the clothing in proximity to the wound. If the muzzle of the firearm is in actual contact with the surface of the body, the entrance-wound will be freely lacerated and ecchymosed in addition to being burnt. Smokeless powders cause neither blackening nor tattooing of the skin.

If a gun loaded with **small shot** is fired into the body when close to it, a somewhat circular aperture, larger than a bullet wound and rather more irregular and contused at the edges, will be produced. There is nothing definite about the exit-wound, because the whole of the original charge scarcely ever leaves the body, if it does, the exit-wound will be larger and still more lacerated than that caused by a bullet, and will, of course, be proportionately larger than its own entrance-wound. Usually the pellets are severally deflected within the body, and do not traverse it *en masse*, the consequence being that the exit-wound may be less than the entrance-wound, very often there is no exit-wound at all. If a gun loaded with shot is fired at the body from a short distance, the surface will be more or less peppered by the pellets, with possibly the production of an irregular wound caused by some which have not spread so widely, at a greater distance, there will be isolated pellet wounds only. It is impossible to assign with accuracy the respective distances at which these various results are produced. Some guns carry much closer than others, the quality of the powder also exercises a considerable influence, and probably the manner of loading.

When both entrance- and exit-wounds are present, a line drawn between them and prolonged on the side of the entrance-wound will enable some idea to be formed as to the situation of the weapon when fired. The information thus obtained is less reliable if the wound has been produced by a spherical bullet than with a conical projectile, because of the greater liability of the former to deflection. When the position of the deceased at the moment the weapon was fired is known, or can be inferred, the spot whence it was fired may be approximately ascertained, a man has been shot dead when writing at a table on to which the upper part of the body fell forwards, in such a case it is not difficult to allot the position whence the weapon was fired. If the projectile has traversed the body the entrance- and exit-wounds must be differentiated. When there is but one wound, the course taken by the projectile within the body may afford a clue as to the relative position of the firearm and the deceased, possible deflection of the bullet being taken into account. Marks produced by the projectile on neighbouring objects, such as furniture, or on the walls of a house, may afford a clue to the direction in which the weapon was fired.

In the case of *Reg v Monson*, Edin, 1893, the prisoner was accused of having murdered a friend named Hambrough, by firing a gun at him whilst they were out shooting, the dead body being found on a "dyke" or turf wall. On three trees in a line with the body pellet-marks were found which, if caused by pellets from the fatal charge, proved that whilst the deceased was standing on the dyke the gun was fired from a horizontal position, this pointed to homicide. The defence alleged that the marks were of earlier date than that of Hambrough's death. The shot struck the head of the deceased obliquely, *en masse*, which indicated a very short range. From experiments, specially made, Littlejohn concluded that the distance of the weapon from the head of the deceased was between three and nineteen feet, probably nine feet. For the defence Hay stated, as the result of experiments made by him, that it was from a few inches to four feet, or within arm's length. The



weapon being charged with *ambricite*—a smokeless explosive—the absence of blackening, or of powder marks about the wound, was without significance. A verdict of not proven was returned.

Serious wounds may be produced by firearms charged with powder only, especially if the substance used as wadding is of a dense nature. If the wadding penetrates the skin it proves that the weapon was discharged within a few feet of the body.

When the bullet remains within the body it is not to be assumed that it was fired from a distance, its momentum being partly spent, for although a rifle bullet under ordinary conditions would traverse any part of the body if discharged into it at short range, it is not so with all revolvers. A long-barrelled rifled revolver would probably send a bullet through the body if fired at short range, but small pocket revolvers for the most part leave the projectile within the body, when such a weapon is fired at the head, in close proximity to it, the bullet remains within the cranium or is imbedded in some part of the skull.

Attention must also be paid to the holes made in the clothing by the passage of a bullet through it. Trevor<sup>1</sup> has described a case in which the fibres of a chemise surrounding an entrance-wound projected outwards in a direction contrary to that which might have been expected. This was due to the gush of blood washing out the fibres, and then to subsequent stiffening of them by the blood. The outer surface of the cloth was slightly charred.

In estimating the risk to life from wounds produced by firearms it is to be remembered that after the immediate results of the injury are recovered from, secondary hæmorrhage and a variety of inflammatory processes of a kind dangerous to life may set in.

It has been asserted that recognition of an assailant in the dark by the light given off from a firearm discharged near to the person assaulted is not possible. Much depends on the relative position of the parties, if the assailant is well within the field of vision of the person at whom he fires, recognition is quite possible, if at the extreme limit, it is doubtful.

## WOUNDS IN THEIR CAUSAL RELATION.

The duty of the medical witness does not end when he has arrived at a conclusion as to the mode of death. In many cases when a dead body is found without any history—death being due to wounding—the question is asked, were the wounds caused by **accidental, homicidal, or suicidal** violence?

This important and, in some instances, formidable question—formidable on account of the obscurity of the indications from which an answer is to be obtained—involves the methodical consideration of a number of criteria which universal experience has formulated as aids to diagnosis, and, what is of no less moment, a keen observation of the smallest details which are special to each case. A wound has to be considered in relation to its **position**, its **nature**—whether incised, contused, etc.—its **direction**, and its **extent**. If the body is examined in the position and at the place in which it was found, circumstances of external relation are to be taken into account—as footmarks, blood-stains, indications of a struggle having taken place, the presence of a weapon in the hand or near the body of the deceased, any change in the position of the body after death, with other matters that present themselves to an observant eye. A minute investigation embracing every perceptible detail should be made—

<sup>1</sup> *Trans. of Med.-Leg. Soc.*, 1913.

note-book in hand—before anything is disturbed, if this precaution is not taken, the unravelling of some important particular may be enhanced in difficulty or even rendered altogether impossible

### THE POSITION OF THE WOUND.

Almost any part of the body may be wounded by a second person—certain parts are inaccessible to the suicide, and certain others are preferentially selected by him. These axioms, for the most part, hold good, but they are not without exceptions. The **front** and more **exposed parts** of the body are usually selected by the **suicide**, the **throat** and **chest** for **incised wounds** and **stabs**, and the **orbit**, **temples**, **mouth**, and **cardiac region** when **firearms** are resorted to. The position of suicidal wounds varies in relative frequency in different countries, in Great Britain one of the commonest modes of committing suicide is by an incised wound of the throat, on the Continent, cut-throats, in comparison with other suicidal wounds, are not nearly so numerous. Stabs in the throat are suspicious of homicide, but instances are not wanting in which suicides have inflicted upon themselves wounds of this description. Incised wounds of the throat, apart from any other injuries, point to either suicide or homicide, and away from accident. Stabs in the back are more likely to be homicidal than suicidal, but, with the exception of the area covered by the scapulæ and the space between them, a man might stab himself in the back were he so disposed. The ordinary suicide selects an ordinary position on which he inflicts wounds for the purpose of putting an end to his life, with lunatics the case is different, they are just as likely to put an end to themselves in some unheard-of way as they are to adopt one of the more usual methods. Little<sup>1</sup> reports the case of a woman, aged thirty-six, recently discharged from an asylum, who, with a blunt table-knife, made a wound in the back of her neck which half severed the head from the trunk, all the tissues were divided as far as the spinal canal, which was opened, the cord just escaping injury—she died on the sixth day after inflicting the wound. In another case,<sup>2</sup> a man was found standing with a large gash in his abdomen, from which he was pulling out his intestines, a coil of the ileum, almost severed, was outside the abdomen, he lived a few hours. It is interesting to note that a well-marked intussusception of the ileum of several days' standing was present.

**Contused** wounds on the **head**, when not due to accident are indicative of **homicide**, but an insane person may inflict such wounds on himself. Smith<sup>3</sup> relates the case of a man who placed himself before a looking-glass, and struck repeated blows on the top of his head with a hammer weighing nearly 3 pounds. An area 3 inches in diameter was divested of scalp, and a fracture of the skull 2 inches in diameter, depressed  $\frac{3}{4}$  inch, was produced, the bones being splintered around. Staples<sup>4</sup> gives an extraordinary instance of self-inflicted injuries to the head which came under his notice, and supplements it with thirteen other cases by various observers. A man drove into his head two stone-chisels, each  $8\frac{1}{2}$  inches long, and  $\frac{3}{4}$  inch in diameter, using for the purpose a wooden mallet weighing  $2\frac{3}{4}$  pounds. One of the chisels was driven through the head from right to left, entering in the right temporal region, and emerging on the left nearly in a direct line, the point projecting  $1\frac{1}{2}$  inches, the head of the chisel being close down to the scalp, the other chisel was driven into the centre of the forehead, penetrating at least  $\frac{1}{2}$  inch into the frontal lobe. After inflicting

<sup>1</sup> *The Lancet*, 1889

<sup>2</sup> *London Med. Recorder*, 1890

<sup>3</sup> *Med Times and Gaz*, 1878

<sup>4</sup> *Journ Amer Med. Assoc*, 1887

the injuries, the man, with the chisels in his head, approached a glazed door, through which he was seen by two persons, he stooped and tried to unlock the door, but did not succeed in doing so. When the door was broken open, he walked a distance of 40 feet with but little aid, and was able to talk. The chisels were withdrawn with considerable difficulty, and he died about five hours afterwards. There is much to be learnt from these cases as regards the immediate effects of severe injuries to the head, in both, repeated blows were struck without producing unconsciousness, and in the second case the patient was not only conscious, but he could talk and walk in spite of the desperate nature of the injuries inflicted on the brain. Such instances are to be remembered by the medical witness when asked as to the possibility of similar wounds being self-inflicted, and also as to the possibility of individuals retaining consciousness and power of locomotion and speech after the receipt of severe injuries to the head, whether self-inflicted or not.

**Suicidal** wounds from blows on the head are usually inflicted within a limited area, and have more or less the same direction, they are generally on the top or front of the head being parts most accessible. The presence of other wounds on the body may be significant—Are they all compatible with suicidal causation, or do they rather resemble homicidal wounds? Indications of resistance, especially on the hands and arms, should be looked for. **Homicidal** head-wounds are often on or towards the occiput, the victim being attacked from behind. If the victim is not rendered insensible by the first blow, he will involuntarily put up his hands to protect his head, and, consequently, the backs of the fingers will probably be bruised. **Accidental** wounds on the head are usually on the vertex when produced by falling head downwards, and partake of the nature of an injury caused by a single blow, which, however, if the body falls from a great height, may be of sufficient force as to smash the vault of the cranium into fragments. Accidental wounds resulting from stones and like objects being projected through the air and striking the head may, of course, be found on any part of it.

Wounds of the **male genital organs**, if not accidental, are for the most part self-inflicted. Ablation of the penis or of the testicles, or even of both, scrotum included, is not an uncommon act on the part of a man labouring under sexual monomania, or under some form of insanity in which the fixed idea prevails that the sexual organs or functions are the cause of his misery, occasionally, wounds of the male genital organs are criminally inflicted out of revenge. Wounds of the **female external genitals** are chiefly accidental, or are due to criminal violence.

Incised or punctured wounds of the **limbs** may result from criminal violence or from accident, in the former case, they are not unfrequently brought about by attempts at self-protection. Occasionally suicides make incisions into the arms and legs, with the object of dividing blood-vessels so as to cause death from hæmorrhage.

## THE NATURE OF THE WOUND.

**Contused** wounds are usually either accidental or homicidal, they are rarely of suicidal origin. **Incised** and **punctured** wounds may be homicidal, suicidal, or accidental, the probability in each case being governed by their position and extent. So far as position goes, the same may be said of wounds produced by **firearms**.

Unusual methods of causing death sometimes betray their homicidal or

suicidal origin, and at others, in the absence of circumstantial evidence, or of that of an eye-witness, they leave the matter in doubt. Of the former class are those exceptional cases in which a man prepares an elaborate apparatus for self-destruction. For example, a man constructed a guillotine in such a way that a suspended axe-blade was liberated after a certain amount of water had run out of a can, in the bottom of which was a hole—the loss of weight caused by the water flowing from the can eventually released a detent by which the axe-blade was held up. An open cavity was prepared, in which a large quantity of ether was exposed immediately under the nose of the suicide. The axe fell in due time and decapitated the constructor of the machine, probably after he had been narcotised, or, possibly, was dead from inhalation of the ether vapour.<sup>1</sup> In a case reported by Leadman,<sup>2</sup> a man committed suicide by placing a dynamite cartridge in his mouth, lighting the fuse and then awaiting the explosion. The soft palate and tongue were torn and mutilated, the teeth broken off, the superior maxillary bones separated and fractured, and the inferior maxilla was broken into about twenty pieces, notwithstanding all this, the skin of the lips and cheeks was intact, the man lived two hours. Other similar cases have occurred. Such a mode of death could only result from suicide or accident, and much more probably the former.

As an instance of death caused in an unusual way, which would give rise to suspicion of homicide in the absence of eye-witnesses, the following, recorded by Stephens,<sup>3</sup> is a striking example.—A man suffering from melancholia (who not long before had been discharged from an asylum), whilst at work forging nails, was seen with a red-hot iron rod, about two feet in length, the cool end of which was against the wall, and the heated end against his belly. One of his fellow workmen gave him a push and made him drop the iron, he said that he should be all right if he was allowed to go on with his work, and he was permitted to do so. Not long after he made the iron white-hot, and succeeded in thrusting it four or five inches into the abdomen, he died the following day.

### THE DIRECTION AND EXTENT OF THE WOUND.

Most men and women are right-handed, and, consequently, incised and punctured wounds suicidally inflicted usually take more or less a definite direction, due to the weapon being wielded with the right hand, but apart from the exceptions which obviously occur in the case of left-handed persons, the indications afforded by the direction of a wound are not to be regarded as absolutely distinctive. It is quite true that the direction of a wound often enables a correct opinion to be formed as to its suicidal or homicidal origin, still, should the direction not agree with that which is held to be characteristic of suicidal wounds, suicide is not, therefore, to be excluded from consideration, unless the position and direction are such as to make it impossible for the wound to have been self-inflicted. Experience teaches the necessity of great caution in applying general rules to special cases, in all doubtful cases allowance must be made for exceptional occurrences.

**Incised wounds** of the **throat** self-inflicted by **right-handed** persons usually run from **left to right** in an oblique direction, the beginning of the cut being at a higher level than its termination. In producing them the blade of the razor or knife is applied to the left side of the throat above the thyroid cartilage

<sup>1</sup> *Boston Med and Surg Journ*, 1880

<sup>2</sup> *Brit Med Journ*, 1881

<sup>3</sup> *Bristol Med-Chir Journ*, 1888

and is drawn obliquely downwards across to the right. Such is the rule, but it has its exceptions. In a case recorded by Mackenzie,<sup>1</sup> a man cut his throat with a razor, the wound was on the right side of the throat, and extended from about the angle of the jaw to nearly the middle line of the neck on a level with the hyoid bone, the direction being from right to left. At its commencement the incision was clean, at its termination it was hacked and irregular. The patient, who was right-handed, afterwards explained that he held the razor (which was blunt) in the right hand, and cut from behind forwards.

Suicidal cut-throat wounds are sometimes made below the thyroid cartilage, such wounds are usually short in length and horizontal in direction, occupying the middle of the throat, between the sterno-mastoids, which frequently escape injury. At other times a clean sweep is made through the whole of the soft structures of the anterior segment of the neck. In suicidal cut-throat wounds the skin is usually the last structure divided—the wound gradually becoming shallower as it reaches its termination, in homicidal wounds of the throat the end of the wound is often under-cut, the skin not being divided as far as the underlying tissues are. Two statements are frequently accepted in relation to extensive wounds of the throat—one is, that after the carotid artery or jugular vein is wounded the person so injured is at once deprived of the power of movement and dies immediately, the other is, that in the case of suicides the incision is never deep enough to implicate the bodies of the vertebrae. Both these statements are shown to be incorrect by a case which happened in the Salford Royal Hospital in 1883. A man was admitted for fractured femur and was placed in the hoist for the purpose of being transferred to the ward. A nurse in an upper storey, who happened to look down into the hoist as it ascended, saw the man take a pocket-knife out of his pocket and apply the blade to his throat, she gave the alarm, the hoist was stopped and a house-surgeon and a porter seized the man to prevent him doing himself further mischief. He had made a large wound, and, notwithstanding all efforts to restrain him, he succeeded in getting his fingers into it and tearing it further open, he died in a few minutes. The author examined the body immediately after death, and found the right carotid artery and jugular vein divided and the body of one of the vertebrae distinctly notched by the blade of the knife. The fractured femur for which the patient was admitted was due to a fall from a height, which, at the time, was supposed to be accidental. It really resulted from an attempt at suicide, and having failed the man took the earliest opportunity of effecting his purpose in another way. It is not unlikely that the cries of the nurse increased his desperation and thus caused him to use unwonted force, hence the injury to the vertebrae.

The extent of the injuries that a suicide may inflict on his throat is exemplified by another case which was admitted into the Salford Royal Hospital, reported by Lord.<sup>2</sup> A man, fifty-eight years old, was brought to the accident room with a large open wound in the throat, he died five minutes after admission, having survived the injury rather more than an hour. Shortly after, his son appeared with something in his hand which he said his father had cut out of his own throat, this turned out to be the entire larynx—the thyroid and cricoid cartilages, together with the first and part of the second ring of the trachea, the whole had been cleanly excised without injuring the large vessels of the neck. Harrison<sup>3</sup> relates a similar case of a woman, aged forty-one, who excised her own larynx, cricoid cartilage, and five rings of the trachea without wounding

<sup>1</sup> *Brit Med Journ*, 1887

<sup>2</sup> *Trans Path Soc Manchester*, 1892

<sup>3</sup> *Brit Med Journ*, 1883

the carotids death quickly followed. In a third case reported by Aezel<sup>1</sup> a woman lived between six and seven hours after she had excised her larynx without injuring the adjacent large vessels. An instance of survival for a time from numerous and severe self-inflicted injuries to the neck and chest is afforded by a case admitted into Middlesex Hospital under Hulke<sup>2</sup>. A man committed suicide by attempting to cut off his head from behind with a shoemaker's knife, as this failed he stabbed himself repeatedly in the chest and finally cut his throat. Crossing the nape of the neck were three deep, jagged, incised wounds, a jagged incised wound crossed the front of the throat from the posterior border of the left sterno-mastoid to the middle of the right sterno-mastoid severing the depressors of the hyoid bone and cutting out a portion of the thyroid cartilage. On the front of the left side of the chest were four stabs, two of which penetrated the pleural sacs. The man survived a week.

**Homicidal** incised wounds of the throat, when inflicted by a right-handed man facing his victim, are from right to left, and are usually more horizontal than suicidal throat-wounds. If the assailant stands behind the victim the wound may closely resemble one of suicidal origin the position and movement of the hand and arm being very like that of a person who inflicts a wound on his own throat. In such a case the incision will be from left to right, and will probably sever the whole of the soft structures down to the vertebra, one of which may be nicked. Very deep and extensive division of the soft structures in front of the throat, especially when associated with nicking of a vertebra is regarded as an indicative of homicide, the indication for the most part holds good, but, as already stated, a like condition may exceptionally be met with in suicidal cut-throats.

**Suicidal** stabs of the chest are usually on the left side in the case of right-handed men, and they take a downward and inward direction. If there is more than one such wound, all will generally be found within a circumscribed area. Multiple stab wounds of the chest of **homicidal** causation are usually distributed over a wider area, and are more horizontal in direction, they may be from below upwards, which is rarely the case in suicidal stabs. The occurrence of several stab wounds on the front of the chest, more than one of which may be sufficient to cause speedy death, does not necessarily contra-indicate suicide. An instructive case of this kind is related by Newnham<sup>3</sup>. A man was found dead with the right hand clenched tightly and a small knife lying in front of him on the floor. On the front of the chest, one inch to the inner side of, and three-quarters of an inch above, the left nipple, were five small wounds, transverse in direction, each about three-quarters of an inch long by a quarter of an inch wide. Just to the inner side of the nipple was another wound half an inch in length, and about one inch below the nipple was a small wound also about half an inch in length. The direction of the six wounds first named was downwards and slightly inwards, all corresponded externally to the third intercostal space, and they penetrated the thoracic wall in the fourth interspace. On the left side of the pericardium was a transverse wound one inch in length, the left ventricle was penetrated by two transverse wounds each three-quarters of an inch long, and the heart was wounded in three other places. All the wounds were of suicidal origin. The multiplicity of wounds in this case might be regarded as indicative of homicide, but against this view is the extremely limited area they occupied, more than one plunge of the blade, which produced a separate external wound, coincided so closely in direction

<sup>1</sup> *Gyógyászat*, 1894

<sup>3</sup> *Brit Med Journ*, 1868

<sup>2</sup> *The Lancet*, 1889

as to make but one large opening in the pericardium, and the two penetrating wounds of the ventricle were only separated by a narrow tongue of the ventricular wall. It would be extremely improbable that a number of wounds homicidally inflicted could be planted so closely together, the struggles of the victim to escape would present a fresh part of the chest wall to the knife each time it fell. The fact that the right hand was tightly clenched, the knife being found on the floor, might, in a doubtful case, have given rise to suspicion. When the weapon that has caused the wounds is found *tightly* grasped in the hand which corresponds with the position and direction of the wounds, the presumption of suicide is strongly corroborated, if it lies loosely in the hand so that it can be lifted away without difficulty, no reliable inference can be drawn, the case may either have been one of suicide, or a second person after inflicting the wounds may have placed the weapon where it was found in order to simulate suicide.

Penetrating wounds of the back are very suggestive of homicide, in exceptional instances they may be due to accident, as when a man falls backwards on a pointed object.

As a means of determining between suicide and homicide when death has resulted from wounds inflicted by cutting instruments, considerable importance is attached to the presence of cuts on the hands and fingers, they are regarded as indications of resistance to homicidal violence, but, exceptionally, they may be met with in suicide. In a case reported by Alexander<sup>1</sup> an officer was found with two deep incised wounds on the front of the abdomen, and one on the back near the spine. There were twenty-six wounds about the left breast, some penetrating the thorax, and both hands were dreadfully mutilated. A sword covered with blood and bent to an angle of 45° was lying beside the patient, who survived several hours, and explained that he had tried to transfix himself by placing the hilt of the sword against the wall and then pressing forward on the point of the blade. On failing he tried a second time, when the blade penetrated the abdomen and impinged on the spine, he withdrew it with great difficulty, his hands being cut in the act of pulling it out. He subsequently attempted to perforate the heart. A somewhat similar case was admitted into University College Hospital under Beck<sup>2</sup>. A man placed the point of a sword-stick against the chest, just below and to the outer side of the left nipple, and drove it in by running against a wall. The blade penetrated eleven and a half inches backwards, slightly downwards and to the right, it was firmly fixed, and was removed with difficulty, having probably pierced one of the vertebræ, at the point of entry one of the ribs was fractured. The man was living when admitted into hospital.

Maschka<sup>3</sup> relates a case which furnished an extreme example of the excess of violence not unfrequently resorted to by lunatics in the suicidal act. A man, aged fifty-one, after being in an asylum for two months, had so far recovered as to be entrusted with a knife for the purpose of cutting an apple. He was afterwards found bleeding profusely from no less than 285 punctured wounds, of which 200 were on the left half of the chest, 50 on the inner side of the left forearm, and 28 on the inner side of the right forearm, the left radial and ulnar arteries were divided. Six of the chest wounds penetrated the thorax, the left lung being compressed by blood collected in the pleural sac. The man survived nearly twenty-four hours, eventually dying from hæmorrhage.

<sup>1</sup> *The Lancet*, 1885

<sup>2</sup> *The Lancet*, 1882

<sup>3</sup> *Prag med Wochenschr*, 1888

### GENERAL CAUSAL INDICATIONS.

Amongst the indications by which an opinion may be formed or strengthened, as to whether death was due to accident, homicide, or suicide, are the **position of the body** when found, the presence on it of **bruises or blood-stains**, the state of the **clothing and surroundings**, the **presence of a weapon**, its position in relation to the body, and, in the case of a weapon other than a firearm, marks of blood on it

The **position of the body** itself may afford a clue to the way in which the wound was caused, especially when considered in relation to the surroundings. When called upon to investigate a suspicious case, careful scrutiny should be made for **foot-marks** on the floor of the room in which the body is found, and also in the adjoining corridors, any passing to and fro should not be permitted until a thorough investigation has been made. If blood has flowed on to the floor, it is probable, in the case of homicide, that the murderer will have trod on it and will have produced marks indicative of his subsequent movements. To avoid error the soles of the feet of the deceased should be examined, as a person, after fatally wounding himself and soiling the floor with blood, might tread in it and then walk about and thus make suspicious footprints, if he has done so, signs of the original hæmorrhage will probably accompany the footprints, as blood will continue to flow from the wounded parts whilst the individual perambulates the chamber. When a murderous outrage has been conducted with great deliberation, the murderer or murderers have been known to remain in the house a considerable time after the death of the victim, and to leave distinct imprints in masses of coagulated blood, these, along with other foot-marks, should be measured, and, if possible, sketched or traced. Marks produced by **blood-stained fingers** on clothing, furniture, or walls should be carefully preserved, as they may be of great value in identifying an assailant (see p 55). For the same reason weapons and other articles which may bear impressions of fingers should be handled as little as possible.

Marks on the dead body or clothing, caused by blood-stained fingers, should be examined with reference to position, a mark on the right side of the body, produced by the fingers and thumb of the left hand, would be a suspicious indication. Finger-stains may exist on parts of the body where they could scarcely be self-produced, as, for instance, between the shoulders. The appearance of blood-spurts on clothing or on furniture has been previously described.

**Bruises** on the body of the deceased may be of great significance, they should especially be sought for on the throat, chest, and arms. Notice should be taken of the presence of marks or indents produced by the finger-nails, they may be so well imprinted as to suggest that the assailant had exceptionally long nails and this may serve as a clue. Too much should not be made of slight ill-defined bruises which may have existed some time before the fatal injury was inflicted.

A **disordered state of the clothing** is suspicious of homicide, but it may be due to frantic movements on the part of a suicide. If the body is in bed, clad in night attire, disarrangement of the clothing is very easily produced, and would only be of significance if sufficiently marked as to suggest that a struggle had taken place. On the other hand, if the bed-clothes are exceptionally straight, they may have been readjusted by the murderer to obliterate previous disorder, in this case the straightening will probably be overdone. Lunatics, before committing suicide, have been known to throw all the furniture in the room into disorder, giving rise to the appearance of a struggle having taken



place, the fury of the insane person may be so maniacal as to cause him to completely wreck the contents of the room before putting an end to his life. Excess of damage to furniture, with probably the presence of a single fatal wound on the victim, would be sufficient to clear up the case, a lesser degree of disorder would be more suspicious, but when a madman begins he usually makes a full end.

Stabs or other similar wounds inflicted through the clothing should be carefully inspected, and the wounds on the body compared with the cuts through the clothing before the latter is taken off. This is especially necessary when the body is fully clothed.

A knife tightly gripped in the hand of the deceased has already been pointed out as an indication of suicide, more frequently it is found lying near the body. When an investigation takes place on the spot, the position of the weapon in relation to the body should be noted before anything is disturbed. In exceptional cases of suicide the knife or razor with the blade closed had been found by the side of the body, in one instance the razor was found in the pocket of the deceased, such an unusual disposal of the weapon probably resulted from persistence of an habitual action which customarily follows the use of a pocket-knife- the folding blade of the razor suggesting the act. When death has been caused by stabs or incised wounds, the absence of a knife or other likely weapon is suggestive of homicide. Sometimes wounds are homicidally inflicted with one weapon and another is left near the body in order to suggest suicide. The implement left is almost always one belonging to the deceased, and it has happened that the only available one was much too small to have produced the injuries found on the body- for instance, a clean sweep of all the soft structures of the neck down to the vertebræ, such as might be made with a carving-knife, could not well be made with a small-bladed pen-knife.

The amount of **blood** found on the **weapon** varies, and may be quite insignificant, even when no attempt has been made by wiping or washing it to remove all trace, thus a long-bladed knife rapidly plunged in and withdrawn may show slight indications of blood, although a large vessel has been divided. This is partly due to quick withdrawal before the bleeding begins, and partly to the blade being clasped by the skin and thus wiped as it emerges, it is still more likely to be free from blood if the wound is inflicted through the clothing. In such cases the blade presents the appearance of having received a thin coating of red lacquer, the actual tint produced being yellow rather than red. A short-bladed knife will probably show more blood near to, and on the handle. Directions have been previously given for examining blood-stained instruments. All that is necessary to say in addition is that, before dissolving off the stain, the blade of the instrument should be scrupulously examined for hairs, etc., with the low power of a microscope.

### THE CAUSAL RELATION OF WOUNDS PRODUCED BY FIREARMS.

With regard to wounds inflicted with **firearms** there are certain special indications which aid in determining whether death resulted from **accident**, **homicide**, or **suicide**. Unless smokeless powder has been used, the chief of these is the presence of particles of powder in the wound, and of general blackening in its vicinity, or of one hand. Blackening of the wound merely shows that

the weapon was fired close to the body, blackening of the hand tends to prove that the wound was self-inflicted, either accidentally or suicidally. Singeing of the hair, scorching of the skin or of the clothing, the character and size of the wound, all yield evidence as to whether the weapon was fired close to or at some distance from the body, the bullet, if found, may give a clue as to the kind and bore of the firearm. The wadding used in charging a shot-gun has been found in the wound, and, being composed of a piece of paper, has been traced by the printing or writing on it to its original owner, at the present day the employment of breech-loading firearms, which necessitates the use of cartridges, lessens the chance of discovery in this way of the person who has discharged the weapon. In all cases any projectile or other substance in the wound should be preserved.

Blackening of the wound being evidence of close proximity of the muzzle of the firearm to the body at the moment of discharge, is absence of blackening (ordinary gunpowder being used) proof of the contrary? When a firearm is discharged at the body within one or two feet, some amount of blackening is almost invariably produced, but exceptional cases are recorded where no blackening of the wound was present, although the firearm was held in the hand of the victim. A case is recorded by Casper-Liman (*l.c.*) in which the dead body of a man was found shot through the heart. There was no blackening, neither round the wound, on the neck, nor on the face, the case, however, was undoubtedly suicidal. Hubbard<sup>1</sup> relates a very interesting case of this kind with some experiments made in relation to it. A man was seen flourishing a revolver, he then sat down under some bushes, a few minutes after a shot was heard, and his body was found with a bullet-hole in the centre of the forehead. The wound was not in the least powder-marked, but all the evidence, except want of blackening, pointed exclusively to suicide. The four remaining charges in the revolver were fired at targets covered with white chamois-leather at distances of 3, 8, 18, and 30 inches respectively, and in every case the target was blackened. It appears from these cases that suicidal death from firearms is not invariably accompanied by blackening of the skin in the neighbourhood of the wound, and therefore, that in suspected homicide by firearms absence of blackening is not inconsistent with the weapon having been fired close to the body—at a distance equal to that from which a man might fire a revolver at himself. The experiments with the targets seem to contradict this, as it would be impossible for anyone to fire a revolver at his own forehead at a greater distance than 30 inches, the teaching of actual cases, however, must be accepted.

Blackening of the hand is not a necessary result of firing a revolver, and, therefore, it may be absent in cases of suicide by means of such a weapon.

With smokeless powder there is no burning of the skin or scorching, and the tattooing is much less evident. Some smokeless powders, for example cordite, cause definite blackening, but not scorching.

It is important for the medical jurist to bear in mind that after a man has inflicted a series of potentially fatal gunshot injuries on himself, he may be able to discharge the weapon a second time into his body. Harvey<sup>2</sup> records the case of a man who discharged a sporting gun loaded with small shot into his left side, the seventh and eighth ribs were smashed, and a wound about four inches below the nipple was produced, which extended about six inches downwards in the direction of the stomach and transverse colon. Notwithstanding this severe injury the man placed another cartridge in the gun and fired it into his mouth. The roof of the mouth and the left side of the face were

<sup>1</sup> *New York Med Journ*, 1887

<sup>2</sup> *Brit Med Journ*, 1895

blown away, the brain was disorganised and the cranial bones were broken into twelve or fourteen pieces, which were found lying loose in the cranial cavity, and yet the scalp remained intact. Revolver-wounds of the head or of the heart are by no means invariably fatal at the moment. It would be natural to infer that a man who had projected a bullet through his heart would be quite unable to fire a second shot through his brain, or, having lodged a bullet in the brain, to shoot himself through the heart. Both of these events, however, have happened, not in one solitary instance only, but in a considerable number of cases. Want of knowledge as to the possibility of such occurrences has led medical witnesses to regard the occurrence of two such wounds, *ipso facto*, as proof of homicide. In a case recorded by Hubbard,<sup>1</sup> a man was found dead in his own barn, his revolver, which contained five chambers (three of which had been discharged), lay within reach of his right hand. There were three bullet-wounds on the body, two of them in the cardiac region - the scorched and powder-burnt clothing indicating very short range, the third was in the right temple, and penetrated the brain at least four inches, this wound was also powder-burnt and was evidently inflicted last. One of the bullets fired at the heart missed it, the other entered the cavity of the right ventricle. This was an undoubted case of suicide. Agnew<sup>2</sup> quotes a number of cases of a similar nature. A student shot himself in the head, walked along a passage to his bedroom, and then shot himself in the heart. A policeman, in the presence of witnesses, shot himself through the head, and then fired a second shot into his chest, the first ball entered the right temple, and was found within the cranial vault on the opposite side, the second ball entered the right side of the heart, death from internal hæmorrhage occurring in about five minutes. A boy, aged nineteen, inflicted with a revolver four wounds on his own person. The first bullet entered the forehead, and, after taking a circuitous route, lodged about the middle of the left temporal lobe, the second bullet passed through the sternum and cut through the left ventricle of the heart, on a level with the mitral valve, a third shot entered the abdomen, and the fourth penetrated the neck, death ensued from hæmorrhage into the pericardium.

Wounds from firearms on the **back** of the body are suspicious of homicide, it is not impossible, however, for them to be self-inflicted either accidentally or, under special circumstances, suicidally. A wound in the back has been accidentally caused by a sportsman dragging after him through a hedge a loaded gun with the muzzle pointing towards him. A gun resting insecurely against some object may be accidentally disturbed and discharged into the back of a bystander as it is falling. Exceptionally, suicides have shot themselves through the back of the head. Haberd<sup>3</sup> examined the body of a man who shot himself through the head in the presence of a number of people in a café, it was found that the bullet had entered exactly at the meeting of the sagittal and the lambdoidal sutures. A man at the moment of pointing a revolver at his heart may be discovered, and in the attempt to wrest the weapon from him, it may be accidentally directed towards his own back, and fired while still grasped in his hand. There is obviously a limit to the angle at which a revolver can be presented to the back whilst in the hand of the suicide, but it is to be borne in mind that, during a desperate struggle, a direction of the barrel quite unattainable by voluntary effort may be accomplished with the unpremeditated assistance of a second person. In such cases, the position of the entrance-

<sup>1</sup> *New York Med Journ*, 1887

<sup>2</sup> *Medical News*, 1887

<sup>3</sup> *Mittheilungen a d Instt f ger Med Wien*, 1891

wound is of less importance than the direction taken by the bullet, allowance being made for deflection caused by its striking a mass of bone, such as the body of a vertebra. If the entrance-wound of an injury suicidally inflicted is situated towards the middle of the back, and the course taken by the bullet is nearly in the postero-anterior line, the muzzle of the revolver must have been close to the body, and there would consequently be blackening bruising, and, probably, scorching of the surrounding surface.

In some cases of suicide with a revolver the weapon is found tightly gripped in the hand after death, this affords as conclusive proof of suicide as any evidence short of that of an eye-witness. More frequently the weapon will be found lying on the ground close to the body. If the revolver lies loosely in the hand the case may be one of suicide or of homicide, in the latter case, the weapon has been placed in the hand of the deceased in order to divert suspicion from the murderer. When a man accidentally shoots himself fatally whilst examining a revolver the weapon falls from his hand, as it is but loosely held at the moment it is discharged, the grip with which a revolver is retained in the hand after death is a prolongation of that due to powerful emotion—with which it was held during life, this is wanting in accidental shooting. Sometimes the weapon cannot be found, although this is suspicious, it is by no means conclusive of homicide, a suicide, after shooting himself, occasionally throws the weapon away, and if this takes place in the open air the revolver may escape discovery, or a passer-by may pick it up without any suspicion that its dead owner is lying a few feet away.

When more than one wound is found on a dead body it may be of importance to determine which was last inflicted, this is not difficult if only one of the wounds is of such a nature as to be speedily followed by death. In some instances there may be two wounds, each of which would be likely to cause immediate death, as, for example, bullet-wounds of the heart and brain. The illustrations already given of double wounding of this description show that sometimes the heart and sometimes the head is first wounded. The course taken by the bullets and the amount of damage they have inflicted on the respective organs may possibly admit of the formation of an opinion, but, unless the indications are well marked, it is advisable not to make a definite statement. The opinion which has been often expressed that immediate insensibility necessarily occurs after bullet-wounds of the brain is not in accordance with facts, it is quite true that immediate loss of consciousness *usually* follows the passage of a bullet through the brain, but exceptions are sufficiently numerous to forbid an unqualified assertion that this necessarily took place in the case of a dead body found thus wounded. In the greater number of the cases above cited the head was wounded before the heart, but they are not to be accepted as conclusive precedents.

### THE CAUSES OF DEATH FROM WOUNDS.

When death results from wounding, medical evidence will be required to prove the **cause of death**, and the relation between it and the injuries found on the body. In a great number of cases the cause of death and its relation to the injuries is patent, in some the cause of death is obscure in others, it is obvious, but its relation to the injuries is difficult to trace. The causes of death may be divided into those which are (a) **immediate**, and those which are (b) **remote**.

(a) **IMMEDIATE CAUSES OF DEATH.**

The immediate causes of death from wounds comprise - **Hæmorrhage, shock, and injury of a vital organ.**

**Hæmorrhage** is a frequent cause of death from wounding. It may occur either rapidly from a wound in a large vessel, or slowly and continuously from division of a number of small vessels. The rate of flow influences the result produced by the loss of a given quantity of blood, an amount of blood, which, in a few seconds, flows from a wound in a large artery and causes immediate death from syncope, may be sustained with impunity if spread over a longer time. The extremes of age, or the presence of debilitating disease, diminish the power of resistance to hæmorrhage, whilst certain diseases and constitutional conditions, as purpura hæmorrhagica and hæmophilia, strongly predispose to it. **Internal hæmorrhage** is frequently a cause of death in perforating wounds of the cavities of the body. In some situations, extravasation of a small quantity of blood is sufficient to cause death from mechanical disturbance, as, for example, in the brain or the pericardium, or from asphyxia, when the trachea is wounded and blood is drawn in through the wound.

**Shock** may cause death either as the result of a **single injury**, or from the **sum of a number of injuries** no single one of which would be fatal. Illustrative of the **first** mode of production are those cases in which death immediately follows a fall from a height, a violent blow on the head, chest, or abdomen, without profuse hæmorrhage occurring either externally or internally. Violent blows on the abdomen, by paralysing the splanchnics, and producing extreme dilatation of the vascular area supplied by them, may deprive the nervous centres of their proper supply of blood, and thus cause death from syncope. In other cases, blows on the chest or belly may cause reflex paralysis of the heart. (See section on Injuries of Chest and Abdomen.)

Illustrative of the **second** mode in which fatal shock may be produced, are cases of severe flogging, examination of the body after death may reveal signs of multiple bruising, but not of any wound or other manifestation of a fatal kind. In a vigorous, healthy person done to death in this way, a very considerable amount of bruising may be reasonably expected to be visible, but in weakly children the post-mortem appearances of injury may be very slight. Such cases occasion the medical witnesses much anxiety, as the cause of death is purely inferential, there is no physical condition to which it can be attributed, and, therefore, it is clear that a positive assertion as to the cause of death is out of place, the witness will have to content himself with stating that, in the absence of other causes, death is referable to shock. He must be prepared for a searching cross-examination, and it is better candidly to avow from the first the impossibility of determining the cause of death with certainty, than to have that statement dragged out of him after an apparent attempt to conceal it. All the organs and tissues of the body should be carefully examined for pathological changes, the defence always being that death resulted from some latent disease.

Severe injury to **organs necessary** for the continuance of **life** is a cause of death sufficiently evident to require no comment. The extensive injuries to the brain or the thoracic viscera which follow explosions, falls from high buildings, railway accidents, and the like, serve to illustrate this mode of death.

Occasionally, examination after death which has immediately followed some act of violence reveals the presence of a pathological condition to which death is due. Hæmorrhage due to advanced degeneration of the cerebral

vessels, to the presence in them of a small aneurism, or to rupture of a thoracic or abdominal aneurism, may occur immediately after the infliction of an injury. Pre-existing cerebral disease may be found after death alleged to be due to violence. A boy was taken ill the day after receiving a slight blow on the head, and died in a few days. The person who had struck the blow was arrested and charged with having caused the boy's death. At the necropsy, old-standing disease of the middle ear was discovered, with an abscess in the cerebellum, also of some standing, the accusation, of course, fell to the ground.

In cases in which death is due to a pathological condition which existed at the time the violence was inflicted the question may be asked. Was death accelerated by the violence? For example, a man with aortic regurgitation, immediately after a tussle with some one drops down dead, there are no signs of violence, nor is there any doubt that disease of the heart was the cause of death. It is well known that both physical exertion and mental excitement are likely to be suddenly fatal in such cases, and usually the accused is exonerated. It is not, however, necessary that external wounding should take place in order to establish a charge of manslaughter, much depends on the nature of the encounter. In the absence of malice or of intention to terrify, the accused would probably receive the full benefit of his inculpableness, but a man who frightened a boy to death by "personating" a ghost was held to be guilty of manslaughter. Death from pure psychical shock without predisposing lesion is very rare. Maschka relates one such case. Two women were in the midst of a furious verbal encounter when one of them seized a besom and made as though she would strike the other, without any blow being delivered the woman who was menaced fell down dead. At the necropsy not the least sign of mechanical injury was found, neither externally nor internally, nothing but hyperæmia of the lungs, and a full right heart, with no other abnormal condition, death being probably due to heart-paralysis caused by excessive mental perturbation. Templeman<sup>1</sup> records the case of a man, aged forty-three, who died immediately after becoming extremely excited, and without being subjected to any physical violence. At the necropsy, sixteen hours after death, both sides of the heart, especially the right, contained a quantity of fluid blood, the heart substance, the valves, and the coronary arteries were free from disease, and no pathological lesion was found to account for death.

### (b) REMOTE CAUSES OF DEATH.

According to the common law of England, a person who inflicts an injury on anyone shall not be held answerable if death is not caused until after the lapse of a year and a day, within this period, if death results from the injury, the assailant may be tried and punished. It is an exceptional occurrence for death due to an injury to be so long delayed, in the ordinary course of events it takes place within a few weeks or months.

The **remote** causes of death from wounds comprise **inflammation, septic processes, and exhaustion**. At a period still further removed, death may result from a traumatic lesion, which for a prolonged interval is either unattended by symptoms, or symptoms may be present during the whole time. An example of the first type of injury is found in rupture of the diaphragm, from the immediate effects of which the patient recovers, but after an interval of several months, during which there may be an entire absence of symptoms, some of

<sup>1</sup> *Edin. Med. Journ.*, 1893

the abdominal viscera pass through the opening and the patient dies of diaphragmatic hernia. The second type is illustrated by injury to the lumbar spinal cord, the patient being paraplegic from the time he sustained the injury, death from bed-sores and general exhaustion taking place after a prolonged interval.

Wounds of a certain kind, or affecting certain organs or tissues, are specially liable to be followed by secondary pathological processes. Severe crushes, or violent stretching of the limbs, may cause rupture of the main artery without any external wound, gangrene being the result. Wounds of the scalp are frequently followed by erysipelas, such wounds are often sustained by intoxicated individuals in brawls, and in the event of death taking place, the defence usually is that the erysipelas was due to drink. Even should this be the case, it does not exculpate the aggressor, but it may be the means of diminishing the punishment allotted to him. Wounds of the brain are sometimes followed by hyperæmia, or inflammation of one or of both lungs, which occasionally is of a peculiarly isolated kind, perhaps one lobe is gorged with blood, and the rest of the organ contains the normal or even less than the normal amount, but it may be œdenatous, the appearance being very like that of one type of *vagus-pneumonia*. It is supposed that the condition arises partly from laboured respiration caused by disturbance of the *vagus-nucleus*, and partly from dilatation of the pulmonary vessels due to paralysis of the *vaso-motor* nerves following the injury to the brain. Another factor may be present—paralysis of the laryngeal muscles with consequent non-closure of the glottis during swallowing, this leads to the introduction of saliva and particles of food into the air-passages, and, as the reflex irritability of the larynx is lessened or abolished, no adequate attempt is made to get rid of the foreign elements by coughing, the result is the occurrence of broncho-pneumonia.

**Septic processes** may cause death within a short time after receipt of an injury, or they may partake of a more chronic character and gradually exhaust the strength by the formation of multiple depots of pus. A patient who has been the subject of homicidal wounding may be received into the ward of a hospital in which an outbreak of septicæmia or erysipelas occurs, and may succumb to one or other of these diseases. The original wound may not have been of a kind dangerous to life, and it will be consequently urged in defence that the accused ought not to be held responsible for a result which in the ordinary course of events would not have happened. The principle laid down by the law is that when a man criminally wounds any person he must abide by the full consequences of the act. It may be true that the said person would not have died had he not been exposed to an infective disease whilst in the hospital, but his presence there was due to the criminal violence to which he was subjected, and therefore the accused is answerable for his death.

**Tetanus** is another disease of an infective nature which may entirely alter the prognosis of a case. A slight wound, equally with a compound fracture, may be followed by tetanus. If death results from tetanus after a wound of a limb it may be stated in defence that had the limb been amputated the injured person would have recovered. Apart from the uncertainty of the benefit which might have resulted from the operation—the benefit being solely hypothetical—it is sufficient if the patient declined to have it performed, the refusal does not lessen the responsibility of the accused. The incubation period of tetanus is usually from one to twenty days, in the tropics the disease has been known to develop in a few hours after the infliction of a wound, but in temperate climes it does not usually supervene until from the fourth to the tenth day. Purves

Stuart<sup>1</sup> states that tetanus bacilli may remain latent in the tissues following a wound for months or even years until further operative treatment or trauma liberates the spores which have been shut up in scar tissue or encysted around some foreign body, and produces a long-delayed local tetanus or even a generalised tetanus-toxæmia. The incubation stage is not unfrequently discussed in medico-legal cases, in relation to which of two injuries caused death from tetanus. For example, a boy had his right foot injured by a horse, in a week he returned to school, and two days after he complained of stiffness in the neck, which he attributed to a thrashing he had received the day before from the schoolmaster, he died from tetanus in seventeen days. At the inquest, medical evidence was given to show that on account of the shortness of the period which intervened, the beating was probably not the cause of the tetanus; the disease showed itself within twenty-four hours after the administration of the correction, whereas the injury to the foot, both from its nature and from the interval that elapsed before tetanus developed—ten days—was an exceedingly probable cause.<sup>2</sup> The incubation period of tetanus may be so far prolonged that doubts may arise as to whether an attack is, or is not, due to a particular injury inflicted some time before. It is exceptional for tetanus to set in after the third week from the occurrence of the injury, if it does recovery is more likely to take place than when the disease develops earlier. Huntly<sup>3</sup> records the case of a boy, aged thirteen, who sustained a lacerated wound of the right foot followed by tetanus, which developed on the twenty-third day after the accident, recovery taking place. Of 367 cases of tetanus which occurred during the American war,<sup>4</sup> in seven the disease developed from the twenty-sixth to the thirtieth day, and in twenty-three, not till the thirtieth day. In one instance it is stated that the disease did not make its appearance until seven months after the injury was sustained.

Of 1,092 cases analysed by E. W. Hill, in 17.49 per cent. the incubation period was from one to five days, and in 55.06 per cent. from five to ten days. In only 8 cases was the incubation period as long as twenty days.<sup>5</sup>

**Idiopathic** tetanus is of more than doubtful existence. It is now generally accepted that traumatic infection is a necessary factor in the causation of tetanus, and that in cases of so-called idiopathic tetanus some slight causal injury has been overlooked.

### SELF-INFLICTED WOUNDS FEIGNING HOMICIDAL VIOLENCE.

Such wounds are inflicted with the following motives — revenge, or a desire to injure the reputation of some one who is accused by the wounded person of having inflicted the wounds, to avert suspicion, as when a man who has charge of money or valuables appropriates them and pretends that he has been robbed with violence, to obtain sympathy or notoriety, the subjects usually being hysterical girls or women, or, after committing murder, to make it appear that the wounds were inflicted by a third person who was guilty of the murder, or (it may be alleged) by the murdered person himself, who having attacked and wounded the survivor, was killed by him in self-defence.

Self-inflicted wounds of this type are almost invariably either stabs or incised wounds, and are limited to parts of the body accessible to the individual himself.

<sup>1</sup> *The Diagnosis of Nervous Diseases*, 1920

<sup>2</sup> *Brit. Med. Journ.*, 1892

<sup>3</sup> *The Lancet*, 1885

<sup>4</sup> *The Med. and Surg. History of the War*, Part III, Surg., 1883

<sup>5</sup> Quoted by Osler, *Principles and Practice of Medicine*, 1919



They are frequently numerous, and if so, are spread widely apart, they rarely penetrate below the cutis, and if they do, they are not on those parts of the body where wounds are popularly supposed to be mortal, in the two last-named characteristics they differ from suicidal wounds. The hands are seldom wounded, whereas in genuine homicidal violence this frequently happens from attempts at self-defence. A further and most important indication is afforded by a careful comparison between the wounds on the body and the cuts through the clothing by which it was covered, or alleged to have been covered, at the time the wounds were inflicted. In all cases a *direct comparison is essential*, the clothes must be placed on the body as they were worn at the time. It frequently happens that some of the wounds have been made whilst the surface was uncovered, the designedly corresponding cuts through the clothing having been made separately. In this case discrepancy is inevitable there will be cuts through the clothing without corresponding wounds on the body, and, what is more significant, there will be wounds on the body covered by clothing which is intact. In one case of this kind there was an incised wound of the arm, but the cut in the sleeve of the coat, which was supposed to correspond, did not divide the lining, it only went through the outer cloth. Besides discrepancy between the cuts through the clothing and the wounds on the body, various portions of clothing may themselves yield contradictory results. Stabs inflicted through the ordinary clothing must pierce more than one layer of it, often several, before the skin is reached. It is almost impossible to arrange the various folds of clothing, such as are presented by the coat, waistcoat, shirt, and under-vest, when off the body, so that the cuts through them will coincide when they are on it.

The medical witness should further carefully examine and compare each article of clothing at the parts perforated, with regard to blood-stains. If the suspected person possesses an ordinary degree of cunning he will not fail to smear the edges of the cuts with blood, this he will probably do to each garment separately. The result will be that in some instances two contiguous layers of cloth will be found unequally stained, the outer layer, possibly, being more freely stained than the one that underlies it.

It is said that individuals who have attempted suicide and have failed to kill themselves, out of a feeling of shame have accused some unknown person of homicidal wounding. Such a case would form a remarkable exception to the rule which obtains in cases of self-inflicted wounds with fraudulent intent, inasmuch as the idea of giving a false explanation of the origin of the wound (probably only one) having been conceived after it was made, the wound would not possess the characteristics which have been described, on the contrary, it would most likely be in a dangerous situation, and might have been produced by a firearm.

## CHAPTER XXV

**PROFESSIONAL PRIVILEGES, OBLIGATIONS AND RESPONSIBILITIES.**

A REGISTERED medical man, who undertakes the treatment of the sick, receives certain privileges in the exercise of his profession, and incurs various obligations the non-fulfilment of which may subject him to severe pains and penalties. Some of these obligations are imposed by statute, others are obligations at common law. In addition there are duties, such as that of taking a dying declaration, the sanction for which is custom and moral obligation rather than enforcement by law.

The **privileges** which the law confers upon a registered medical man place him in regard to certain matters in a position better than that of the non-registered practitioner. He is empowered, for example, to sign certificates required by various Acts of Parliament, such as death certificates, certificates under the Lunacy Acts, Education Acts, etc. He can give certificates excusing witnesses or jurymen from attending in court on the ground of ill health, and these will ordinarily be accepted in courts of law. He is exempt from the necessity of serving on a jury, he is entitled to sue for his fees, and he can hold public appointments.

**Registration.**—It must be pointed out that both the privileges and responsibilities of the medical practitioner are acquired by the act of registration. A man may possess the highest medical and surgical qualifications obtainable, but unless he has been registered by the General Medical Council his position is no better than that of any unqualified person. A medical man, therefore, who has taken a registrable qualification and intends to practise, should see that his name is placed upon the register without delay. If for any reason, such as prolonged absence from the country, or failure to receive communications from the Registrar at the time of revision of the list, his name has been withdrawn, he should, before recommencing practice, take the steps necessary to have it replaced.

In many Acts of Parliament it will be found that the words "duly qualified" medical practitioner are used. By Section 34 of the Medical Act these are construed to mean a person registered under that Act.

**The General Medical Council.**—This body, created by the Medical Act of 1858, consists of persons appointed by the Crown through the Privy Council, persons appointed by the various Licensing Bodies and Universities in the British Isles, and a certain number of representatives elected directly by the registered medical practitioners in the British Isles. It maintains the Register of medical practitioners and possesses disciplinary powers over the profession. Other duties are to exercise some control over courses of study prescribed for medical students, to see that an efficient standard of knowledge and skill is required for the passing of qualifying examinations, and to issue the list of drugs and medicines with methods of preparation known as the British Pharmacopœia.

It is the exercise of the disciplinary functions which concerns us here. Section 29 of the Medical Act runs as follows: "If any registered Medical Practitioner shall be convicted in England or Ireland of any felony or misdemeanour, or in Scotland of any crime or offence, or shall after due inquiry be judged by the General Council to have been guilty of infamous conduct in any professional respect, the General Council may, if they see fit, direct the Registrar to erase the name of such Medical Practitioner from the Register." The definition of 'infamous conduct in a professional respect' was laid down by Lord Justice Lopez in *Allinson v General Medical Council* (63 L Q B 534). He said: "If it is shown that a medical man, in the pursuit of his profession, has done something with respect to it which would be reasonably regarded as disgraceful or dishonourable by his professional brethren of good repute and competency, then it is open to the Council to say that he has been guilty of infamous conduct in a professional sense."

A succession of cases which have been heard and decided by the Council have now practically determined the offences which are regarded as falling within the meaning of the words quoted. The more important of these are:

A conviction for a criminal offence, unless of a very trivial character.

Immoral relations between doctor and patient where the professional relationship exists.

Signing untrue, misleading, or improper medical certificates, notifications, reports, or kindred documents.

On certain matters the Council has issued the following pronouncements—

**Unqualified Assistants and Covering.**—The employment by any registered practitioner in connection with his professional practice of an assistant who is not duly qualified or registered, and the permitting of such unqualified person to attend, treat, or perform operations upon patients in respect of matters requiring professional discretion or skill, is in the opinion of the Council in its nature fraudulent and dangerous to the public health, and any registered practitioner who shall be shown to have so employed an unqualified assistant is liable to have his name erased from the Register.

"Any registered practitioner who by his presence, countenance, advice, assistance or co-operation, knowingly enables an unqualified or unregistered person, whether described as an assistant or otherwise, to attend, treat, or perform any operation upon a patient in respect of any matter requiring professional discretion or skill, to issue or procure the issue of any certificate, notification, report, or other document of a kindred character, or otherwise to engage in professional practice as if the said person were duly qualified and registered, is liable on proof of the facts to have his name erased from the Register."

"The foregoing do not apply so as to restrict the proper training and instruction of *bona fide* students, or the legitimate employment of dressers, midwives, dispensers, surgery attendants, and skilled mechanics, under the immediate personal supervision of a registered practitioner."

**Sale of Poisons.**—The employment, for his own profit and under cover of his own qualifications, by any registered practitioner who keeps a medical hall, open shop, or other place in which scheduled poisons or preparations containing scheduled poisons are sold to the public, of assistants who are left in charge but are not legally qualified to sell scheduled poisons to the public, is in the opinion of the Council a practice professionally discreditable and fraught with danger to the public, and any registered practitioner who is proved to have so offended will be liable to have his name erased from the Register.

**“ Association with Unqualified Persons.**—Any registered practitioner who, either by administering anæsthetics or otherwise, assists an unqualified person to attend, treat, or perform an operation upon any other person, in respect of matters requiring professional discretion or skill, will be liable on proof of the facts to have his name erased from the Register

**“ Advertising and Canvassing.**—The practices of (a) advertising by a registered medical practitioner with a view to his own gain, particularly if depreciatory of other practitioners, or of sanctioning such advertising, of (b) employing or sanctioning the employment of agents or canvassers for the purpose of procuring patients, and of (c) associating with or accepting employment under any Association which practices canvassing or advertising for the purpose of procuring patients, are in the opinion of the Council contrary to the public interest and discreditable to the profession of medicine, and any registered medical practitioner resorting to any of such practices renders himself liable on proof of the facts to have his name erased from the Medical Register

**“ Association with Uncertified Women practising as Midwives.**—Whereas it is provided in the *Midwives Act*, 1902, and in the *Midwives (Scotland) Act*, 1915, that ‘ no woman shall habitually and for gain attend women in child-birth otherwise than under the direction of a qualified medical practitioner unless she be certified under this Act ’, and whereas it has been made to appear to the General Medical Council that certain qualified medical practitioners have, from time to time, by their countenance or assistance or by issuing certificates, notifications, or other documents of a kindred character, knowingly enabled uncertified women, on pretence that such women were under their direction, to attend women in child-birth, contrary to the law, and whereas such conduct is, in the opinion of the Council, discreditable to the profession of medicine, and calculated to defeat the purpose of the Statutes made in the public interest for the protection of mothers and infants, notice is hereby given that any registered practitioner who is proved to have so offended will be liable to have his name erased from the Medical Register ”

In less flagrant instances the Council may postpone further consideration of the case for a time in order to give the offender an opportunity of discontinuing the practice complained of

There is no appeal from a decision of the General Medical Council, when given after due inquiry and without malice

### THE OBLIGATION TO EXERCISE REASONABLE SKILL AND CARE.

The law requires a medical practitioner to exercise **reasonable skill and care** in the treatment of his patients. Failure to do so constitutes the offence of **malapraxis**. If a patient suffers in consequence of negligent treatment, the medical practitioner responsible may have a civil action brought against him and may be mulct in heavy damages. If gross carelessness or negligence has been displayed, a criminal prosecution may follow. The important question in relation to this subject is—What degree of skill or of care is necessary to exonerate a practitioner from the charge of malapraxis? The question only permits of a relative answer, there is no arbitrary standard by means of which a sufficiency of skill can be estimated, and the same may be said of the amount of care bestowed. The qualifying expression, “reasonable,” admits of considerable latitude of interpretation, a greater degree of skill would be expected

from a town physician or surgeon of high standing than from a country practitioner. The opportunity for calling the higher skill into action, however, may not occur, the treatment of a simple fracture of the arm can be carried out by a country practitioner equally well as by a hospital surgeon. In such a case any flagrant error in treatment would be blamable in one as much as in the other. Any medical practitioner may make a mistake in diagnosis or in treatment, but that is not sufficient to establish a charge of malapraxis, it must also be proved that he did not bring a reasonable amount of skill and attention to bear.

It will be observed that malapraxis may be due to one or both of two causes—want of reasonable skill, and want of reasonable attention and care. A surgeon may perform an operation with the requisite skill, but if, for its success, great care and attention are afterwards necessary, and these are not bestowed, he lays himself open to an action for damages. The alleged subsequent neglect may be capable of such an explanation as to make it a debatable point, or it may be of such a nature as to leave no doubt of its existence. In a severe case of epistaxis, for example, a surgeon may skilfully plug the nares, and then leave the plug until it becomes putrid, and septic infection results. If accused of negligence, he might say that the bleeding was so profuse that he did not think himself justified in removing the plug at an earlier period. This is a feasible explanation, and the most that could be made of the case would be, that there had been an error of judgment, but such a mistake would scarcely be held sufficient for damages. On the other hand, a degree of negligence may be exhibited by a medical practitioner in omitting to relieve urgent symptoms which an average knowledge of his profession would inform him ought to be treated, that renders him liable to legal proceedings, as, for example, leaving a patient with a full bladder so long unrelieved that serious consequences ensue. Even where the treatment itself is of a kind proper to be undertaken, failure to observe precautions sufficient to prevent untoward secondary effects may constitute negligence. For instance, in 1904 damages were awarded against a medical man who used X-rays and high frequency currents in a case of malignant disease, whereby the patient suffered excessive dermatitis and burning of the skin.

Apart from absence of reasonable skill and care, a medical man may render himself liable to an action for damages if he adopts some entirely new mode of treatment which partakes more or less of the nature of an experiment, the result being unfavourable. It is seldom that this can occur, since improvements in treatment are of gradual development, and, although a new method may be quite different from what was formerly adopted, still it will either have sufficient relation to the older treatment to deprive it of the nature of a pure experiment, or it will be based on physiological or pathological facts, by appeal to which the new departure can be justified. It would be unreasonable that medical science should be restricted in its advances as regards treatment, this is so generally appreciated at the present time that new methods of treatment are welcomed and willingly submitted to by the public, although they are perfectly cognisant of their tentative nature. Whenever, in a particular case, a medical practitioner contemplates the adoption of a method of treatment that differs to a material degree from the methods which are recognised by the profession at large, he should take his patient into his confidence, explain the nature and object of the treatment, and, before putting it in force, should obtain his freely given consent. If the patient, by reason of tender years or from any other cause, is incapable of appreciating the difficulty, the consent

of the nearest relative should be obtained after all the facts of the case have been laid before him

From time to time actions are brought against surgeons for having unskilfully treated a fracture of the arm or leg, the patient further emphasising his view of the matter by declining to pay the bill. In such cases, a curious compromise is occasionally effected by the Court. Although there may be no evidence of want of reasonable skill or care, yet if the limb displays an altered appearance, a reduction is made in the amount payable to the surgeon, notwithstanding the fact that the judgment on the main issue is in his favour.

The following is an instance of negligence so gross as to amount to crime—

At the Warwick Assizes, in 1875, a medical practitioner was sentenced to six months' imprisonment for the manslaughter of a married woman whom he had attended in her confinement. After delivering the child with forceps he was observed to cut off something and throw it into a privy. This was proved to be fifteen feet of intestine. At the autopsy the vagina was found to have been ruptured, and part of the ileum and colon was missing, probably it had been pulled down in the first instance under the impression that it was the cord.

If, whilst in the fulfilment of his professional duties, a medical practitioner is culpably negligent, as, for example, when performing an operation, or whilst attending a woman in labour, the plea that he was under the influence of a drug taken for a legitimate purpose will not exonerate him.

A medical man (*Reg. v. Wright*, C.C.C., Nov. 1895) was indicted for having caused the death of a pregnant woman whilst delivering her with the forceps. At first it was alleged that at the time the prisoner was under the influence of drink, subsequently, it was explained that he had taken a large dose of chloral hydrate some hours before being called to the labour. In summing up, Wright J. said that it might be proper for a man to stupefy himself with chloral, or similar medicine, for the treatment of disease, but that if he did so he must not, while in that state, attend a case of childbirth. The prisoner was found guilty, and was sentenced to three months' imprisonment, the judge stating that if the maltreatment had resulted from alcoholic intoxication a much severer sentence would have been passed. The onerous responsibility of medical practitioners in case of malapraxis, in which death results, is illustrated by the announcement made by the counsel for the defence in this case—that a civil action claiming damages from his client had been commenced by the husband of the deceased.

**Responsibility in a Case of Criminal Wounding.** The position of a medical man in charge of a case of criminal wounding likely to prove fatal is one of great responsibility, if the wounded person dies the treatment he has adopted may be severely criticised. The object of the counsel for the defence is to transfer the blame as far as possible from his client to some other person, and the medical attendant is usually made the scapegoat, hence the necessity of treating such a case throughout with the full knowledge of what is likely to occur. Operative interference must only be resorted to when absolutely necessary. Before any operation is performed a consultation with one or more additional surgeons should be held. The same precaution should be taken if there appears to be any ground for considering an operation needful, since at the trial a medical man may be blamed, on the one hand for having performed an operation which caused the death of the wounded person, or, on the other, for neglecting to have recourse to an operation which would probably have saved his life. If the medical man brings to bear a reasonable amount of care and skill, the culpability of the accused is not affected whatever the issue of the treatment may be. Even if, through mistaken diagnosis, an operation is performed after due consideration and with a full conviction on the part of the surgeon of its

absolute necessity for the purpose of saving life, and on examination after death it is found that the operation was unnecessary, and hastened or even caused death, still the person who inflicted the original wound is criminally responsible. The exonerating conditions, so far as medical men are concerned, are—that the original wound is likely to prove fatal, that the operation in the opinion of the surgeon is absolutely necessary to save life, and that a reasonable degree of skill and care were brought to bear in the performance of the operation.

If a wound is not originally of a fatal character, and death of the wounded person is caused by unskilful or negligent treatment, the accused is relieved of responsibility. If, for example, a popliteal aneurism resulted from an act of criminal violence, and it was mistaken for an abscess and laid open with the knife, death resulting from hæmorrhage, the person who inflicted the original injury would not be held responsible for the death. The law in requiring that a reasonable degree of skill should be exhibited in the treatment of these cases, means such skill as is supposed to be possessed by every duly qualified practitioner. In remote country places it is not to be expected that the services of operating surgeons of great experience are to be obtained as in large towns, nor does the law demand such a high standard of efficiency. A general practitioner residing in a large town, who has charge of a critical case of criminal wounding, for his own sake should avail himself of the assistance of a hospital surgeon of experience in order to meet the almost inevitable criticism to which the treatment of the case will be subjected in the event of a fatal issue, a practitioner in the country must bring his utmost skill and attention to bear, he has then done all that the law requires.

Death caused by chloroform or other anæsthetic administered for the purpose of performing an operation would be judged by the same rules as if it were due to the operation itself—Was the anæsthetic necessary, and was it administered with reasonable skill and care?

#### **Responsibility of Medical Man in Relation to Hospital Authorities and Nurses.**

— When a medical man has been assisted in his treatment of a patient by a nurse, or has left her to carry out instructions in his absence, and negligence is alleged, the question arises as to who is the party responsible. Complete discussion of this question would involve reciting legal arguments which would be out of place here, but certain broad principles may be indicated.

The general rule of law is that a principal is responsible for the acts of his agent, *qui facit per alium, facit per se*. The particular applications of this rule depend upon the circumstances. During the performance of an operation, the operating surgeon is in supreme control, and is, therefore, responsible for acts of negligence on the part of the nurses, who are, for the time being, in law, his servants. The hospital authority does not incur responsibility provided it has exercised due care in the selection of the nurses.

The legal relations between a hospital authority, surgeon, and nurse were clearly stated in the case of *Hulley v Governors of St Bartholomew's Hospital* (1909, 2 K B 820). In this case the plaintiff sought damages for injuries due to negligence of the defendants at the hospital. The plaintiff entered the hospital in order to be examined under an anæsthetic by a consulting surgeon attached to the hospital. For the purposes of the examination he was placed upon an operating table in such a position that his arms hung over the sides of the table. His left arm was burned by coming in contact with a hot-water tin, which projected from beneath the table, and his right arm was bruised by the pressure of some person against it. Traumatic neuritis and paralysis of both

arms followed. At the original trial, Mr Justice Grantham had refused to leave the question of negligence to the jury on the ground that if there had been negligence, the operation was under the control of the operating surgeon, for whose action the governors were not responsible, nor were they liable for the negligence of their staff even if proved. This view was upheld by the Court of Appeal. In the course of a written judgment, Lord Justice Farwell said: "It is impossible to contend that the surgeon or the acting assisting surgeon or the acting house-surgeon, or the administrator of anæsthetics, or any of them, were servants of the defendants in the proper sense of the word: they are all professional men employed by the defendants to exercise their profession to the best of their abilities, according to their own discretion, but in exercising it they are in no way under the orders or bound to obey the directions of the defendants. The only duty undertaken by the defendants is to use due care and skill in selecting their medical staff.

The nurses and carriers stand on a different footing. If and so long as they are bound to obey the orders of the defendants, it may well be that they are their servants, but as soon as the door of the theatre or operating room has closed upon them for the purpose of an operation (in which term I include examination by the surgeon) they cease to be under the orders of the defendants, and are at the disposal and under the sole orders of the operating surgeon until the whole operation has been completely finished, the surgeon is for the time being supreme, and the defendants cannot interfere with or gainsay his orders. This is well understood, and is indeed essential to the success of operations. No surgeon would undertake the responsibility of operations if his orders and directions were subject to the control of or interference by the governing body. The nurses and carriers, therefore, assisting at an operation cease for the time being to be the servants of the defendant, inasmuch as they take their orders during that period from the operating surgeon alone, and not from the hospital authorities. The contract of the hospital is not to nurse during the operation, but to supply nurses and others in whose selection they have taken due care."

An interesting case illustrating the principles laid down above was that of *Byrne v Thorne*, which was heard in the King's Bench in 1904. The defendant performed an abdominal operation upon the plaintiff which was successful, but subsequently the patient experienced further discomfort. She consulted another doctor, who advised that a second operation should be performed, and at this it was discovered that the plaintiff was suffering from an abscess caused by a mattress sponge which had not been removed after the first operation. She rapidly recovered, and then brought an action for damages for negligence. For the defendant it was argued that it was not part of the duty of the operating surgeon to count the sponges, as his attention should be concentrated on the operation and condition of the patient. It was justifiable to leave the counting to the nurse. In this case two large mattress sponges and twenty-four swab sponges were provided. After the operation the defendant twice asked the nurse if the sponges were correct, and received satisfactory replies. Mr Justice Bruce left the following questions to the jury: (1) Was the defendant guilty of want of due and reasonable care in counting or superintending the counting of the sponges? (2) Was the nurse employed by the defendant as assistant during the operation? (3) Was the nurse negligent in counting the sponges? (4) Was the counting of the sponges a vital part of the operation which the defendant undertook to see properly performed? (5) Was the nurse under the control of the defendant during the operation? The jury answered all these questions in the affirmative and returned a verdict



for the plaintiff, awarding one farthing damages. On reconsideration of the damages by direction of the judge, the jury increased them to £25. In entering judgment, the judge remarked that he was glad the jury had returned a verdict which cast no reflection on the defendant's skill as a surgeon.

Where the nurse is not acting under the immediate supervision of the medical man responsibility does not ordinarily arise, thus in *Perionowsky v Freeman and Holmes* in 1866 (4 F & F 977) it was sought to make two surgeons at St George's Hospital responsible for injuries due to scalding a patient who had been placed by nurses in a bath heated to an excessively high temperature. It was proved that the bath had been ordered by the defendants, but actually administered by the nurses. Lord Cockburn, Chief Justice, directed the jury that the defendants would not be liable for the negligence of the nurses, unless near enough to be aware of and able to prevent it. Persons who went as patients into hospitals were not to be treated with negligence, but on the other hand medical gentlemen who gave their services gratuitously were not to be made liable for negligence for which they were not personally responsible. Our great hospitals supported by alms and voluntary contributions, could not be carried on if they had to engage a staff of medical men sufficient to attend to all the minor incidents or details of medical or surgical operations. It was indispensable that such matters should be left to nurses who were necessarily familiar with them, and such was the ordinary and usual course of hospital practice. The jury returned a verdict for the defendants.

**Malapraxis by Unregistered Practitioners.**—A considerable number of unqualified persons are practising medicine in this country, and as medical practitioners not infrequently meet with cases which have been negligently or unskillfully treated by such persons, it is desirable to indicate their degree of responsibility for malapraxis in the eyes of the law. If an unqualified person represents himself to be legally registered, or allows himself to be so regarded, a more serious view will be taken of his negligence. At the Central Criminal Court in 1882 an unqualified practitioner was found guilty of manslaughter for having allowed a patient with a full bladder to remain unrelieved so long that although an operation was subsequently performed by a surgeon, he died from extravasation of urine. It has frequently been remarked that quacks escape the result of their unskilled treatment when duly qualified medical men would be held criminally responsible. To a certain extent this is so, as a consequence of the rule already stated—That the degree of skill regarded by the law as reasonable is held to vary with the circumstances of each case. Gross want of knowledge may render even a quack responsible, but if the ignorance displayed is of a qualified nature, it may be held that less knowledge and skill are to be expected from a man who has not received a medical training than from one who has, and, therefore, that the culpability incurred is proportionally less.

When criminal proceedings are taken for wrongful treatment which does not cause the death of the patient, more must be proved than is required to obtain a verdict in a civil action. The following case illustrates this. Some so called Indian oculists, who had obtained money by performing certain barbarous operations on the eyes of a number of ignorant people, were indicted before the Central Criminal Court in 1893, for alleged conspiracy to obtain money by false pretences, from divers persons, with intent to defraud. Expert evidence showed that the operations were useless, cruel, antiquated, and barbarous, and that the operators were absolutely incapable of diagnosing diseases of the eye. To convict on this it was necessary to prove not only that the pretence was false, but also that the accused knew it to be false. In summing up the Common Serjeant said to the jury — "If you think that these men deliberately performed these operations with the full knowledge that that which they were doing was useless, unnecessary, and cruel, as the skilled surgeons tell you, you cannot resist the conclusion that the intention they had was to

defraud If you think that this is not established, then they are entitled to be set free " The jury returned a verdict of not guilty Under like conditions, an action for *malapraxis* in a civil court would probably have resulted in heavy damages against the defendants

**Operations and Consent.** -An operation cannot be performed upon a responsible person without the consent of that person In the case of a child, permission should be obtained from the parents or guardian Where a person is not in a condition to give consent, as for instance one brought into hospital in a state of coma, a surgeon is justified in taking such steps as appear necessary to save life, but even in this case, unless delay would be dangerous, permission should be obtained from the nearest relative Before undertaking an operation the surgeon should always explain fully its nature and possible after-effects If, as in some abdominal operations, it is impossible to determine beforehand the extent of the interference which may be necessary, the surgeon should obtain a written permission to use his discretion in doing whatever appears to be in the best interests of the patient An action which was brought against a surgeon in 1896, well illustrates the inconvenience a medical man may be subjected to if he fails to take this precaution The plaintiff, a hospital nurse, who suffered from symptoms indicating disease of the ovaries, went into a home in order that an operation might be performed by the defendant, a consulting surgeon According to her statement, she had instructed the surgeon to leave one ovary even if diseased, but both were removed, and she was in consequence sterile The defendant stated that the decision was left to his discretion, that he found both ovaries diseased, and that their removal was necessary in order to prolong life He also expressed the opinion that the plaintiff was sterile before the operation owing to the disease A verdict was found for the defendant, but the case was taken to the Court of Appeal, and when the verdict was there upheld, the plaintiff endeavoured to go to the House of Lords, but was refused permission to sue *in forma pauperis*

### OBLIGATION OF SECRECY.

The position which medical men occupy as depositaries of secrets, acquired in their professional capacity, concerning the physical or mental condition and the doings or misdoings of their patients, may become one of great delicacy and difficulty It may be stated as a general proposition that a secret which is acquired by a medical practitioner from a patient, whether orally or by means of an examination of the person, remains the secret of the patient, it is imparted to the practitioner for a specific purpose on the implied understanding that the knowledge thus obtained will be used solely in the patient's interest But there are obviously many cases in which it is the duty of the practitioner to inform others of the information he has gained If, for example, a patient is suffering from severe melancholia, the medical attendant is undoubtedly right in informing the responsible relatives of the risk of suicide, in order that proper precautions may be taken In some cases of serious illness it is wiser to inform the relatives only of the gravity of the condition Another group of instances where professional information must be revealed are the various notifications of infectious diseases, births, causes of deaths, etc., which the practitioner is bound by statute to make to a local authority These obligations must be fulfilled even though the effect is highly prejudicial to the interests of the patient or others Thus if a doctor finds a case of scarlet fever in a school, he is bound to inform the medical officer of health for the district, even though

this may lead to closing of the school and serious loss and inconvenience to many persons. A notification of birth cannot be withheld on the ground that the mother is unmarried. A certificate of death must state the true cause of death, though, in this instance, strict obedience to the law is not always observed. The existing system under which the certificate must be handed to the nearest relative, unfortunately, leads practitioners to use terminology concealing the real nature of the disease in cases where it reflects upon the character of the deceased. There is reason to believe that, in consequence, the Registrar-General's statistics of death from alcoholism, venereal disease, abortion, etc., are seriously in error. In so acting, the practitioner, however good his motive, is not fulfilling his legal obligations. Finally in the courts of law a medical man must, if the judge so direct, answer any question that is put to him (see p. 7).

The cases which present real difficulty and demand care on the part of the practitioner if he is to avoid subsequent trouble, are, for the most part, those involving questions of morality. A family doctor, for example, is asked by a mistress to examine a domestic servant whom she believes to be pregnant. In this case the medical man must not only have the consent of the girl to make the examination, but must explain fully the object of the proceeding, and receive permission from her to make known to the mistress the results thereof. If he does not take this course, he renders himself liable to actions at law.

**Privileged Communications.**—Cases arise from time to time in which, irrespective of consent, a medical man may feel that he is under a moral obligation to impart his knowledge to others for their own protection. By making a statement which is defamatory of another, he may render himself liable to an action for libel, unless it can be shown that the communication was *privileged*. 'A privileged communication may be defined as a communication which on its face would be defamatory and actionable, but is prevented from being so by reason of circumstances rebutting the existence of malice. It may exist where any person having an interest to protect, or having a legal moral or social duty to perform, makes a communication to another (such other having a corresponding interest or duty), in protection of his interest or in performance of his duty, here, although the communication may contain matter that would ordinarily be actionable, yet it is not actionable if the communication is fairly and honestly made in *bona fide* belief of its truth and without gross exaggeration.'<sup>1</sup> It is often a difficult matter to determine whether a moral or social duty exists such as to make the communication privileged, and it is then for the judge to decide whether the principle can be applied in the particular case. But where ordinarily the communication would be privileged, it is open to the plaintiff to show that the defendant was, in fact, actuated by malice, such as anger, or with a knowledge that the statement was untrue, or with a reckless disregard of whether it was true or not.

The question of privilege may arise when a medical man is treating a patient for syphilis and considers it his duty to warn others—a wife or nurse, for example—of the risk of infection. The position is a difficult one, and few cases bearing upon it have come into court, probably because the persons chiefly concerned are usually willing to adopt the reasonable and proper precautions advised by the doctor. The following cases illustrate the operation of privilege in this matter—

*Guy v Green* (Leeds Assizes, 1903). A medical man was called in to attend a barmaid at an hotel, and was requested by the manager to see her. He examined her in the presence

<sup>1</sup> Indermaur, *Principles of Common Law*

of the housekeeper and, after making the examination, he informed the housekeeper and the employer that the girl was suffering from venereal disease, and advised that she should go home. Subsequently he made a similar statement to a man who represented himself to be the husband of the girl, though this representation was afterwards shown to be false. It was also alleged that he had made a similar communication to another barmaid, but this the defendant denied. At the trial, the judge upheld the plea of privilege as applying to the communications to the housekeeper, to the manager, and to the man who had represented himself to be the husband, but left it to the jury to decide upon the evidence as to the communication to the other barmaid. The jury found a verdict for the plaintiff with £75 damages.

*Still v Morris* (Q B D 1900) In this case the plaintiff, a fireman in the service of the Metropolitan Fire Brigade, claimed damages against a doctor for negligence and libel. The plaintiff alleged that the defendant had treated him for enlarged glands in the groin and a rash on the body, and had negligently and unskilfully diagnosed that he was suffering from syphilis, that he had given a certificate to that effect to the Metropolitan Fire Brigade, and that in consequence the plaintiff was compelled to leave his employment. For the defence it was stated that the defendant was the District Medical Officer of the London County Council, that as such it was his duty to attend members of the Fire Brigade within the district, and to furnish certain reports and certificates, that he had diagnosed that the plaintiff was suffering from syphilis, and had certified accordingly, and that the certificate was a privileged communication. It was further stated that the sixth clause in the conditions of service ran, "Every fireman is liable to immediate dismissal for unfitness, negligence, or misconduct." The judge held that the communication was privileged, and dismissed the case with costs against the plaintiff.

**Professional Secrecy in Relation to Criminal Matters.** In the course of his professional work a medical man sometimes becomes aware that crime has been committed, either by recognising the indications of criminal violence during his examination of the patient, or by an actual confession being made to him. But while it is the duty of a good citizen to render reasonable assistance in the prevention and detection of crime, the practitioner must always remember that the information gained at the bedside is confidential, and only strong reasons will justify him in revealing it to others. It is not part of his duty to act as a detective and give information to the police whenever he comes across crime. Yet at the same time he must use reasonable discretion, and in a serious case he may be justified in subordinating the immediate personal obligation to public duty. For instance, if a murder or grave assault has been committed, and a person corresponding to the published description of the assailant comes to a practitioner for treatment, the latter should in this case inform the police of the fact. On the other hand, in a case of criminal abortion, provided the woman is recovering, there is no need for him to take steps which would lead to her appearance in a public court and perhaps seriously damage her reputation.

In this connection the statement made by Mr Justice Hawkins in the case of *Kutson v Playfair* is important. He was speaking of the divulgence of crime by medical men, and said "It was also said by the medical witnesses that if in the course of professional practice they came across a case which indicated either that a crime had been committed, or was about to be committed, that under these circumstances they were bound to divulge it. To whom? To the Public Prosecutor! If a poor, wretched woman committed an offence for the purpose of getting rid of that with which she was pregnant, and of saving her character, her reputation, and, it might be, her very means of livelihood, and if a doctor was called in to assist her— not in procuring abortion, for that in itself was a crime— but called in for the purpose of giving her medical advice—how she might be cured so as to go forth about her business—he doubted very, very, very much whether he would be justified in going forth and saying to the Public Prosecutor, 'I have been attending a poor young woman who has been trying to procure abortion with the assistance of her sister. She is

now pretty well, and is getting better, and in the course of a few days she will be out again, but I think I ought to put you on to the woman' To his mind a thing like that would be monstrous cruelty He did not know what the jury's view would be, he spoke only of his own Therefore, when it was said that there was a general rule existing in the medical profession, that whensoever they saw in the course of their medical attendance, that a crime had been committed, or was about to be committed, they were in all cases to go off to the Public Prosecutor, he was bound to say that it was not a rule which met with his approbation, and he hoped it would not meet with the approbation of anyone else There might be cases when it was the obvious duty of a medical man to speak out In a case of murder, for instance A man might come with a wound which it might be supposed had been inflicted on him in the course of a deadly scuffle It would be a monstrous thing if the medical man might screen him, and try to hide the wound which might be the means of connecting the man with a serious crime "

The right course of action for a medical practitioner to take in a case of criminal abortion when serious symptoms are not present is indicated by this statement Of course, if the woman is dangerously ill, the practitioner must give information to a magistrate or to the police in order that her depositions may be taken, and if death is imminent, he must take a dying declaration himself in the manner already described In every case where a medical man knows or suspects that abortion has been criminally procured, he should for his own protection call in a brother practitioner, or at least inform some responsible person of the circumstances

**The Medical Man and "Undue Influence."**—Owing to the confidential relation which exists between the two, a medical man has exceptional opportunities for influencing the mind of a patient It behoves him, therefore, to be extremely cautious when there is any possibility of the allegation being made that he has influenced the patient's mind in a direction to serve his own interests This applies particularly to gifts made by the patient to the doctor during life, or legacies left in a will It is an established principle of law that where confidential relations exist between two persons of such a character that one is likely to acquire habitual influence over the mind of the other, and in the absence of independent advice, gifts are made to that one by the other, undue influence will be presumed from the mere fact of the relations between the parties The onus of proving that such was not the case rests upon the person who received the gifts The strictness with which this rule is enforced is shown by the following case —

*Radcliffe v Price* (Chan Div 1902) The plaintiffs, the executors of a lady who died in March, 1900, sought to recover from her medical attendant gifts amounting to £800, which had been given to him at various periods in 1899 and 1900 The defendant had attended the deceased for eleven years No charge of fraud or improper conduct was made, but it was claimed that in view of the relations between the parties and the absence of independent advice obtained by the donor, presumption of influence existed and that the gifts were not valid It was shown that the sum given was only a small fraction of the total property, which was stated to amount to £90,000, and that the deceased had provided for her husband, and had left legacies to relatives, and donations for a museum, public garden, and various charities Nevertheless, the defendant was required to refund the money and pay the costs of the action Mr Justice Swinfen Eady, in giving judgment, said, "It has been laid down that the relation of patients and physician is a confidential relationship, and where it exists, as it did in this case, the donor must have had competent and independent advice before a gift can be supported."

In view of the legal position, a medical man should make a rule never to

accept a gift from a patient unless he is satisfied that a responsible member of the patient's family has been informed of all the circumstances. When, too, he is aware that a legacy has been left to him in a will, he should see that the fact is brought to the knowledge of the other persons chiefly interested in the disposal of the testator's property. There is no objection to a medical man assisting a person to make a will provided the foregoing principles are observed, but it should be remembered that if he acts as a witness to the testator's signature, any bequest made to him in the will is invalidated.

**Notification of Births.**—The following provisions of the Notification of Births Act 1907, are of importance to medical practitioners—

"(1) In the case of every child born in an area in which this Act is adopted it shall be the duty of the father of the child, if he is actually residing in the house where the birth takes place at the time of its occurrence, and of any person in attendance upon the mother at the time of, or within six hours after, the birth, to give notice in writing of the birth to the medical officer of health of the district in which the child is born in manner provided by this section.

"(2) Notice under this section shall be given by posting a prepaid letter or postcard addressed to the medical officer of health at his office or residence, giving the necessary information of the birth within thirty-six hours after the birth, or by delivering a written notice of the birth at the office or residence of the medical officer within the same time, and the local authority shall supply without charge addressed and stamped postcards containing the form of notice to any medical practitioner or midwife residing or practising in their area, who applies for the same.

"(3) Any person who fails to give notice of a birth in accordance with this section shall be liable on summary conviction to a penalty not exceeding twenty shillings. Provided that a person shall not be liable to penalty under this provision if he satisfies the Court that he had reasonable grounds to believe that notice had been duly given by some other person.

"(5) This section shall apply to any child which has issued forth from its mother after the expiration of the twenty-eighth week of pregnancy, whether alive or dead."

Experience has shown that it is not always safe for the practitioner to rely upon notice being given by another person. To avoid the risk of being summoned he should make a practice of sending the notification himself. In a case where it is difficult to determine whether the twenty-eighth week of pregnancy has been reached or not, it is preferable to err in the direction of giving an unnecessary notice, rather than to refrain from notifying. It should be noticed that the local authority is only authorised to supply stamped postcards. If, as in the case of a still-birth or an illegitimate birth, it is desirable to avoid undue publicity, the cost of postage of a letter must be obtained from another source. The Notification of Births Act is now in force in all areas.

**Notification of Disease.**—In this section only the obligations attaching to medical practitioners as such are considered. For the special duties applying to school medical officers, officers of institutions, etc., books on public health should be consulted. Under the **Infectious Diseases (Notification) Act, 1889**, "Every medical practitioner attending on or called in to visit the patient shall forthwith on becoming aware that the patient is suffering from an infectious disease to which this Act applies, send to the medical officer of health for the district a certificate stating the name of the patient, the situation of the building, and the infectious disease from which, in the opinion of such medical practitioner, the patient is suffering."

The diseases scheduled in the Act are small-pox, cholera, diphtheria, membranous croup, erysipelas, scarlatina or scarlet fever, typhus, typhoid, enteric, relapsing, continued and puerperal fever, but the local sanitary authority has power, with the consent of the Ministry of Health to add other infectious diseases such as measles, chicken-pox, etc. Under orders issued by the Ministry of Health, all cases of tuberculosis, ophthalmia neonatorum, and encephalitis lethargica are now made notifiable. When a local authority declares a disease to be notifiable, notice of each declaration must be sent to every medical practitioner practising within its jurisdiction. The practitioner notifying is entitled to a fee of 2s 6d if the case occurs in his private practice, and of 1s if the case occurs in his practice as medical officer of any public body or institution. Failure to notify renders him liable to a fine of 40s.

A medical practitioner should remember that the obligation to notify an infectious disease still rests upon him if called in to visit a patient, even though he may not be the regular medical attendant of the patient, and is not undertaking the responsibility of treating the case. In a recent instance a doctor was called in to see a maidservant at the house of her employer, and found that she was suffering from scarlet fever. It was decided to send the girl to her home at once. On being summoned for failing to notify, the doctor explained that he did not regard the girl as his patient, and concluded that the notification would be made by her own medical attendant, but this did not save him from being fined and required to pay the costs.

Under the **Factory and Workshops Act, 1901**, "Every medical practitioner attending on or called in to visit a patient whom he believes to be suffering from lead, phosphorus, arsenical or mercurial poisoning, or anthrax, contracted in any factory or workshop, shall (unless the notice required by this subsection has been previously sent) send to the Chief Inspector of Factories at the Home Office, London, a notice stating the name and full postal address of the patient, and the disease from which, in the opinion of the medical practitioner, the patient is suffering, and shall be entitled in respect of every notice in pursuance of this section to a fee of two shillings and sixpence, to be paid as part of the expenses incurred by the Secretary of State in the execution of this Act." Other diseases may be added to the list by special order of the Home Office.

Failure to make the required notification renders the practitioner liable to a fine not exceeding forty shillings.

**Certification of Death.**—Section 20 of the **Births and Deaths Registration Act, 1874**, provides that "In the case of the death of a person attended during his last illness by a registered medical practitioner, that practitioner shall sign and give to some person required by the Act to give information concerning the death, a certificate, stating, to the best of his knowledge and belief, the cause of death." The persons required by the Act to give information concerning the death are the nearest relatives present at the death or in attendance during the last illness and, in default of such relatives, each person present at the death. The certificate of death must be sent by the person giving information to the local Registrar of Births and Deaths. A medical practitioner who refuses or fails to give a certificate of death without reasonable excuse renders himself liable to a fine of forty shillings. As a duplicate may be used for improper purposes, not more than one certificate should be given. It is not legally necessary for a medical man to see the dead body before giving a certificate, but it is highly desirable that he should always do so. A certificate cannot be withheld for the reason that the practitioner's fees have not been paid, nor can a fee be claimed for giving the certificate.

Care should be taken to make the information given in the certificate as accurate and complete as possible, otherwise further correspondence and inquiry may ensue. The local registrar is the first authority to scrutinise the certificate, and if for any reason it appears on the face of it unsatisfactory, he forwards it (or should forward it) to the coroner for the district, who makes inquiries and decides whether an inquest is necessary. Later, when the certificate reaches the Registrar-General, if vague or ambiguous terms have been used, the medical man may be written to and asked to supply further information.

**Refusal of Certificate. Giving Information to Coroner.**—According to the strict rendering of the law a practitioner is justified in giving a certificate even in the case of a death from violence, provided he has attended the deceased in his last illness and knows the cause of death, and there is no statutory obligation upon him to inform the coroner of the circumstances. But if this course were adopted the information would only reach the coroner through the registrar after an interval, and serious delay might occur before a proper investigation into the cause of death was set on foot. It is, therefore, the custom for the medical practitioner to refuse to certify, and to give information himself direct to the coroner, as soon as he becomes aware of the death from causes other than natural, either of one of his patients, or of a person to whom he has been summoned after death. This avoids delay, and it is highly desirable that in the vast majority of cases the custom should be observed.

The particular circumstances under which the certificate should be refused require consideration. Section 3 (1) of the Coroners Act, 1887, provides that "Where a coroner is informed that the dead body of a person is lying within his jurisdiction, and there is reasonable cause to suspect that such person has died either a violent or an unnatural death, or has died a sudden death of which the cause is unknown, or that such person has died in prison, or in such place or under such circumstances as to require an inquest in pursuance of any Act, the coroner, whether the cause of death arose within his jurisdiction or not, shall, as soon as practicable, issue his warrant for summoning not less than twelve nor more than twenty-three good and lawful men to appear before him at a specified time and place, there to inquire as jurors touching the death of such person as aforesaid."

There is no legal definition of a "violent" or an "unnatural" death, and indeed it would not be easy to give a scientific definition of natural death other than death from old age—probably a rare occurrence. Deaths, accidental, suicidal or homicidal, which are violent in the accepted sense of the word, occasion no doubt, and should always be notified to the coroner at once. But difficulty may arise with deaths from causes usually secondary to injury, such as tetanus, septicæmia, or other forms of blood-poisoning. Where there is a history of injury, information should be given. In other cases the practitioner must be guided by the views of the local coroner. It has already been pointed out (*vide p. 3*) that the practice of coroners varies within considerable limits. Some consider it necessary to hold inquests in all deaths from tetanus, others do not do so if there is no history of an injury. Even where septic infection has clearly followed an injury some coroners do not hold inquests if they are satisfied from the preliminary investigation that there is no suspicion of crime or negligence. Strict consistency would, of course, require that every death from bacterial invasion through a breach of continuity of the skin should be regarded as unnatural. Similar uncertainty prevails in the case of deaths under anæsthetics. Some coroners take the view that they are bound to hold inquests in all such cases, others only if there are allegations of negligence. Sometimes



endeavour is made to distinguish between deaths under and deaths from anaesthetics. In all these cases the practitioner should remember that he is not the judge as to whether an inquest should be held or not, and if he has any doubt, he should report the case. Having once learnt the views of the coroner, he will know how to act in future.

Sudden death does not necessarily demand an inquest. If the practitioner is satisfied that death was due to natural causes he is justified in giving a certificate. Such a death would be that of an elderly person who was known to be suffering from serious cardiac lesions and had quite recently been under treatment. If, however, the death occurs in a street or other public place, it is desirable to inform the coroner and withhold the certificate pending his decision.

Perhaps the most difficult class of cases is that in which the symptoms during life have been compatible with disease, but some unusual feature, or possibly the behaviour of an individual, has aroused suspicion of poisoning. There are many poisons which produce effects closely resembling disease, and correct diagnosis from the symptoms alone may be anything but an easy matter. Examples are afforded by the illness of the persons murdered by Kłosowski, described on p. 334. Fortunately these cases are rare, but, when one arises, the medical man is placed in an extremely difficult position. On the one hand, if he refuses to give a certificate, and, at the inquest, his suspicions are proved to be groundless, he will have given the matter an undesirable degree of publicity which may have unfortunate results both for the relatives of the deceased and for himself. On the other hand, if, stifling his doubts, he certifies the death, and fresh facts come to light after the body has been buried, the even more painful proceeding of an exhumation may be ordered. Not a few instances in which this has occurred might be cited. Beyond indicating that the practitioner should not allow himself to be swayed by considerations of self-interest, no general principles can be laid down. Each case must be decided upon its merits after review of all the circumstances of the illness and death.

A medical man is not forbidden by law from making an autopsy, provided he has the consent of the relatives to do so. In many hospitals post-mortem examinations are made as a matter of routine, but in private practice, except under instructions from the coroner, it is rarely advisable to make one. Practically this course is only permissible where doubt is confined absolutely to a question of pathology, and it is desired to clear up a point for the sake of giving an accurate certificate of death. The practitioner should never make a post-mortem examination on his own authority if there is the least suspicion of death having been due to causes other than natural. In view of the possibility of an inquest, nothing should be done which might destroy valuable evidence, or, in a case of poisoning, interfere with organs or material which should be submitted untouched to an expert for analysis. A coroner has no power to order a post-mortem examination and decide on the results thereof whether or not to hold an inquest. Nevertheless, the Editor has met with cases in which a coroner has told a medical man that if he cares to make a post-mortem on his own account and report again later, he (the coroner) will reserve his decision until that has been done. This comes very close to an evasion of the law, but if the practitioner acts upon the suggestion, he must remember that he is accepting entire responsibility, and in the event of adverse comment being subsequently made, he cannot shelter himself behind the authority of the coroner. There are reasons for believing that if coroners had the right, under suitable conditions, of ordering a post-mortem examination without

necessarily holding an inquest, a considerable number of these inquiries would be avoided

An inquest is not rendered unnecessary by the lapse of any interval, however long, between the occurrence of the violence and the death. In 1913 an inquest was held upon a man who had been injured in a railway accident nineteen years before.

In the course of his preliminary investigations before holding an inquest, a coroner will frequently send his officer to a medical man to make various inquiries. In the great majority of cases the practitioner should give freely all the information he possesses, but there is no legal obligation upon him to answer any of the questions, and if for any reason he prefers to reserve his evidence until he is in the witness-box he is strictly within his legal rights in so doing.

## PART II.—INSANITY AND OTHER ABNORMAL MENTAL STATES.

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### CHAPTER XXVI

#### THE FORMS OF MENTAL DISORDER.

To the medical jurist, unsoundness of mind is of importance in relation to several questions, the more important of which are the placing of lunatics under restraint, the responsibility for crime, and testamentary capacity. Each of these questions involves legal principles which will be more fully considered in the succeeding chapter. For the immediate purpose it is sufficient to note that the law has never attempted to lay down any hard and fast definition of insanity, or other abnormal condition. Nor from the clinical point of view is any satisfactory definition possible. Broadly speaking, physicians classify abnormal states into the psychoses—a term which is practically equivalent to insanity and the psychoneuroses and neuroses, which are described as functional in character. In practice, however, border-line cases occur in which the diagnosis may only become clear with the passage of time. As regards the psychoses, various classifications have been proposed, but it must be admitted that there is none which is entirely satisfactory, owing mainly to the fact that in many forms of mental disorder our knowledge of the pathology is still very incomplete. The following classification is based upon that now employed by most text-books of psychiatry. The student must not suppose, however, that the types can always be so clearly differentiated, and in practice overlapping forms are frequently met with.

##### Manic Depressive Psychoses

- (a) Manic types
- (b) States of depression or melancholia
- (c) Alternating, periodic, or circular insanity

##### Dementia Præcox

- (a) Simple dementia præcox
- (b) Hebephrenia
- (c) Catatonia
- (d) Paranoid type

##### Acute Confusional Psychosis

##### Paranoia

##### Epileptic Psychosis

##### General Paralysis of the Insane

##### Toxic Psychoses

##### Psychoses with Somatic Disease

##### Senile Psychoses

Psychoses associated with Reproduction  
Defective Mental Development

- (a) Idiocy
- (b) Imbecility
- (c) "Moral Insanity"

**Diagnosis of Insanity.** - The diagnosis of insanity may or may not be a matter of difficulty. In some psychoses, for instance, general paralysis of the insane, the diagnosis may be assisted by the presence of physical changes, such as the Argyll-Robertson pupil. In many forms also, for example, confusional psychosis, dementia præcox, and manic-depressive types, the diagnosis when the condition is well established is quite clear. Difficulty, however, arises in the early stages of insanity, and in the differentiation of these from certain types of the psychoneuroses. These cases are usually spoken of as "border-line," and frequently the diagnosis does not become clear until they have been under observation for a period, or definite symptoms have appeared with the progress of the disease. The reaction of the patient to treatment is often a valuable indication, and it is safe to say that the modern methods of treatment based upon psychoanalysis have been the means of restoring to normal health persons suffering from psychoneuroses who, not many years ago, would have been regarded as definitely insane and possibly placed under restraint. The student may again, however, be reminded that for medico-legal purposes we are not concerned with the precise differentiation between a psychosis and a psychoneurosis or neurosis. When legal questions are at issue the law is not concerned with the exact diagnosis of the condition, but with the question as to how far the condition has affected the conduct of the individual.

Before proceeding further, it may be useful to define certain terms which are employed in psychiatry—namely, illusion, hallucination, and delusion.

An **illusion** is a false perception of an external impulse, it is objective and is limited to one or more of the senses. It affords no indication of insanity, inasmuch as it can be experienced without participation of the intellect. Sane persons may be the subject of illusions, thus, a man who suddenly awakes from sleep sees what appears to be the outline of a human figure in his dimly lighted bedroom, yet he knows that he is alone in the room and that the door is locked, a few moments' reflection, or a closer inspection of the thing seen, reveals that the impression produced has been caused by a dressing-gown suspended from a toilet-stand or other similar object. This is an example of a visual illusion which is corrected by an appeal to the reasoning faculties, or to the evidence afforded by another sense. The causation was objective—the thing seen was wrongly perceived by the visual centre. No delusion resulted—the error was at once detected and the mind remained uninfluenced.

An **hallucination** is a perception without an external impulse, it is subjective in character and affects one or more of the senses. An hallucination is not necessarily indicative of insanity, inasmuch as it may be rejected by the reasoning faculties, but if of a pronounced nature it is usually accepted by them, and in this case determines a delusion. A man who sees some one watching him through an open window when no one is visible, or hears voices speaking when there is absolute silence, is subject to an hallucination. Less significant hallucinations, such as flashes of light, sparks, or figures having a crenated outline, not unfrequently occur in simple neuroses, such as migraine, they may, however, be precursors of mental disorder. Non-existent animals, such as rats or mice, seen and believed in by patients suffering from delirium

tremens, are delusional hallucinations. In simple insanity hallucinations of hearing are the most common.

A **delusion** is a perversion of the judgment by an erroneous perception or conception, unlike the two preceding terms, a delusion necessarily implies disordered intellect. The delusions of the insane are of a personal nature, in an impersonal way people may labour under delusions without being insane. The perennial sequence of religious and spiritualistic frauds, for example, is the result of perversion of judgment in a certain type of mind, a type that eagerly embraces unorthodox ideas without question, such people are not necessarily insane, but they are liable to become so. In them the reasoning powers are defective without essential loss of perceptivity, they see and hear as other people do, but they form erroneous conceptions with regard to what they see and hear, and from them deduce false conclusions. The delusions of an insane person are not grounded on abstract conceptions solely relating to matters outside his entity, they concern himself, if he has delusions about religion, he is convinced that his soul is irretrievably lost, or that he is the Deity incarnate, if about spiritualism he imagines that he is irresistibly influenced by departed spirits, or is possessed by the Evil One himself. In this way an insane person may be subject to delusions without hallucinations.

Some writers use "illusion" and "hallucinations" as convertible terms, and others use "illusion" in a different sense from that above defined. The word "illusion" is unnecessary to the medical witness and need rarely be used.

Insanity does not develop in a moment, there is a period of ingravescence during which the individual affected gradually deviates from his ordinary mental condition. Although the symptoms evinced during the various stages of insanity are irregular and do not follow any definite order, certain of them are of common occurrence, some of which are present in every case, a knowledge of these symptoms is essential to an early recognition of mental disease.

The onset of insanity may be so insidious that, long before any distinctive indications are manifest, and alteration in temperament is noticeable, the individual is different from what he was formerly, he has lost his equanimity. The patient is subject to unaccountable waves of depression which may alternate with periods of excitement, he becomes unwontedly irritable, and is unable to control his temper under the petty annoyances of every-day life, this instability of temper may be the condition that first rouses suspicion in the minds of his friends that insanity is the cause of the change in disposition. A man who has been moody and reserved in his manner for some time is credited with being overtaxed with business, and is thought to be a little out of sorts, but nothing more, at some trifling contradiction or annoyance he suddenly blazes up into a frenzy of passion, and behaves for the moment so like a madman, that the bystanders are at once impressed with the idea that the balance of his mind is impaired.

A change in the emotions is also commonly shown, by transformation of like into dislike, of love into hate. A man shows unwonted impatience at the remarks addressed to him by his wife, whose opinion he previously valued, and then manifests an absolute antipathy to her, a sentiment totally at variance with their former relations. Loss of interest in objects and pursuits which formerly occupied his attention, desire for solitude, perhaps at first shown by avoidance of social intercourse in a general way, and then, in a more special manner, by seclusion from the family circle, are further evidences of perverted sentiments. At this period, the person affected is often quite capable of brightening up in the presence of strangers, or of friends for whom he has a special

liking, he will even remark that he feels better in company—meaning in the society of those with whom he is not necessarily brought into relation. The capacity to fulfil the duties which devolve upon a merchant or professional man may be equal to the requirements, but the work is done in a perfunctory way without the display of any interest.

Amongst the commoner delusions in the early stages of insanity are the conviction that some one generally a member of the patient's family—has commissioned the police, or a private individual, to act as a spy on the person labouring under the delusion, that there is a conspiracy to ruin him or to deprive him of his rights, that attempts are being made to poison him. The last-named delusion frequently causes those who are afflicted with it to arm themselves with a number of bottles of medicine, or parcels of food, and to seek the advice of a medical man, requesting him to analyse the samples brought for poison. It is rather curious that suspicion is not always directed against those who might be reasonably suspected of criminal motives such as members of the family who would be benefited by the death of the deluded person, but druggists and other shopkeepers who are total strangers to him are often the alleged secret assassins, sometimes the allegation of conspiracy is made to account for this inconsistency.

Delusions originating in the **special senses** are also frequent in the early stages of insanity. A man labouring under such delusions will state that he hears voices making disparaging or defamatory remarks about him, or calling upon him to commit certain actions, he sees people deriding or menacing him in the streets, he states that a friend who has been dead for years, walked past his house and looked at him as he went by, that he can taste poison in his food, that an enemy is influencing him by means of electricity, which causes him to feel strange sensations, or that an offensive odour emanates from his body.

Delusions and hallucinations are always looked upon by the layman as strong evidence of insanity, and, as we shall see in the next chapter, the presence of delusions carries considerable weight when irresponsibility on the ground of insanity is put forward in a criminal charge. Nevertheless, it is now recognised that both hallucinations and delusions may occur in psychoneurotic disorders. Visions of deceased persons may appear in some cases, in others there may be delusions of persecution, and in yet others the patient may identify himself with divine personages, or may believe that he is in direct communication with them. Sometimes voices are heard threatening the patient or commanding him to perform certain acts. That these delusions are not indicative of a psychosis is shown by the fact that under psychoanalysis an explanation for the delusion is found and the symptom may be completely removed. Often, too, the patient who hears voices will admit, when pressed, that he does not believe that they are real.

A hereditary taint can be traced in a large proportion of patients suffering from psychoses. In manic depressive insanity, the percentage, according to different observers, ranges from 30 to 80. The defect in the preceding generation or generations is not always definite insanity, in some cases a history of epilepsy or alcoholism is forthcoming.

The following descriptions of the chief forms of insanity do not purport to be complete from the clinical point of view, but are intended to indicate the chief features in each form which are likely to be of importance from the medico-legal point of view —

## MANIC DEPRESSIVE PSYCHOSES.

Kræpelin has shown that mania and many states of depression described as melancholia should now be regarded as different manifestations of one morbid process, to which he has given the name manic-depressive insanity, and under this term he also includes periodic, alternating, or circular insanity<sup>1</sup> Kræpelin's view has been accepted by most English psychiatrists Experience has shown that there are certain fundamental features common to all these groups Further, the different states pass into each other without recognisable boundaries, and may even replace each other in one and the same case For purposes of description, however, it will be convenient to describe separately manic states and melancholic or depressive states

**Manic States.**—These are divided by Kræpelin into Hypomania, Acute Mania, Delusional Mania, and Delirious Mania

**Hypomania** is the slightest form of manic excitement It corresponds to the French "*folie raisonnée*" or insanity without disorder of intellect The perception and memory of the patients are not disordered, and there may even be increased psychic activity, the patient uttering witty remarks, making startling comparisons, and displaying vividness of imagination Nevertheless, there is a lack of unity in the course of his ideas, and he changes frequently from one subject to another While not suffering from actual delusions, he is apt to have an exaggerated opinion of his own importance, and boasts about his aristocratic acquaintances, high mental capacity, etc, indications which may suggest early general paralysis of the insane For the most part the patient is cheerful and exalted, but he is irritable and liable to violent outbursts of rage over very trivial matters He displays an exaggerated fussiness, and is busily engaged in many schemes, but his capacity for real work is diminished. The condition is of importance from the medico-legal point of view, because the morbid state is very likely to be undiagnosed, and their aggressiveness may lead to the appearance of such persons in court Sometimes they start and carry through law suits with great persistence They may commit offences against decency, and may become addicted to alcohol, the condition then being one of the forms of dipsomania Kræpelin states that sexual excitability is considerably increased, and describes several cases in which this led to improper behaviour An elderly father of a family, who had lived a retired life, began to drink champagne with the girl fencers from a circus, another forced his way into a maid-servant's room, and when caught endeavoured to excuse his conduct on the grounds of "midsummer madness", a married lady in each attack conceived a violent passion for any male person in her vicinity, and finally overwhelmed a man 30 years her junior with fervid declarations of love, a servant girl annoyed a captain in the army with numerous love letters which she signed "your fiancée" Accusations of infidelity may be made, and unreasonable jealousy evinced

**Acute Mania.**—Hypomania passes by insensible degrees into acute mania The onset may be sudden or may have been preceded by headaches, weariness, irritability, and outbursts of anger or states of depression The patient becomes restless and fantastic in his conduct He may dress himself up in an extravagant way or go out of the house in his shirt Kræpelin mentions a case of a female patient who jumped into the carriage of a prince for a joke, and of another who went to church in her ball-dress The patient is effusive, loquacious, and incoherent He is abusive, uses foul language, may destroy all within his

<sup>1</sup> *Manic-Depressive Insanity and Paranoia*, trans. by Barclay, 1921.

reach, tear his clothes to shreds, be filthy in his person, and commit indecent actions. Fugitive delusions are commonly present, and usually of an exalted type. Outbursts of violence with threatening and dangerous assaults may occur.

Appetite and sleep are fitful, and the bowels are constipated. At this stage the mind is as unstable as a fallen leaf on a windy day, it passes from subject to subject with phenomenal rapidity, the succession being apparently devoid of coherence. Not unfrequently a certain sequential order may be recognised in the words or phrases uttered, which depends either on association of past experiences or of the words pronounced, usually it is of a mixed type, a word now calling up a past experience, and now suggesting another word of similar sound or meaning. This state may continue for three or four months with partial remission at intervals. Sleep is irregular, a maniac will pass several consecutive sleepless days and nights, will then sleep for several hours, and recommence his previous behaviour on awakening. The tongue loses some of its coating and the appetite often becomes voracious, notwithstanding which the patient grows thinner. Power of endurance is the most extraordinary feature in cases of mania, the continuous violent movements, shouting, and singing that are kept up night and day surpass anything of the kind that can be accomplished by a sane person.

Acute mania may end in recovery, or it may pass into the **chronic** form, from which recovery rarely takes place, although there may be periods of remission. Chronic mania is more or less associated with dementia, into which state the patient may lapse, or he may do so directly from the acute stage. In other cases mania is succeeded by melancholia. **Chronic mania** is characterised by incoherence and delusions, with occasional exacerbations of an exalted type, which, however, are less active than in acute mania.

**Delusional Mania.**—In this condition delusions and hallucinations are more elaborate and are suggestive of paranoid attacks. The patient believes that he is descended from a royal family, that he possesses great wealth, that he has secret powers, or is in direct communication with God. The picture presented in these cases resembles that of the general paralytic. In acute mania the delusions are transient and fugitive, but in the condition now described they, as a rule, persist for a considerable time, and are recurred to and defended by the patient. Excitement is not a marked feature, but there may be sudden outbreaks of violence or passionate weeping.

**Delirious mania** may develop from ordinary acute mania, or it may begin quite suddenly without premonitory symptoms. It is most common in young persons, especially in those that are of excitable or hysterical temperament. It is sometimes preceded by a period of depression followed by loquaciousness which rapidly passes into a state of acute delirium, if delirious mania develops in this way, it is likely to last longer than when it suddenly bursts forth without antecedent mental disturbance. The patient rapidly becomes confused and disoriented. He mistakes those around him and regards them as persons of high importance. Constantly-changing hallucinations are present. Animals and birds are seen in the room, heaven is opening, the Day of Judgment is come, the house is on fire, the patient is swimming in water. Very little appreciation of the surroundings is shown, but when addressed the patient will look up and obey some simple command or give a childish answer to a question. When the delirium is at its height the patient is violently excited, will attack those who seek to restrain him, will tear his clothes, displaying an utter absence of decency, and will shout and swear or sing until his utterances are absolutely



incoherent In this stage there is absence of sleep, the tongue, from the first foul, becomes more and more so, and then dry and brown, sordes are present on the teeth and lips, and the breath is very offensive The appearance during the periods of lessened excitement resembles that produced by an attack of enterica, but there is little elevation of temperature The attack may consist of periods of frenzied excitement with intervals of comparative quiet, and may be accompanied by hallucinations, it may be only of a few hours' or days' duration, or it may last two or three months The absence of sleep in intractable cases is one of the most marked symptoms, a patient may not sleep for four or five consecutive days and nights

**Diagnosis.**—It is distinguished from **delirium tremens** by absence of the half-frightened manner, the tremor, the hallucinations of rats or serpents on the bed, the furtive glances over the shoulders and under the pillows, the constant purposeless picking at the bedclothes, or grasping at imaginary objects—with the retention usually of sufficient intelligence to answer questions—all of which are indicative of that disease The history of the case will help to distinguish delirious mania from the delirium of a **fever** or an acute **febrile disease**. In a fever, or like bodily complaint, there is a period of illness characterised by elevation of temperature, quickened pulse, and other physical disturbances before the delirium comes on The intensity of the delirium is much less than in delirious mania, although in some diseases, as, for example, in the pneumonia of alcoholic subjects, the delirium is often very violent The absence of any characteristic rash and definite course points to mania rather than to fever From **encephalitis** the diagnosis cannot give rise to much difficulty, in this disease there is severe pain in the head, vomiting, rigors, and a tendency to somnolence, which soon passes into coma, the accompanying delirium is less continued and less violent than that of delirious mania In delirious mania, pain in the head is not a frequent symptom, and there is an absence of vomiting, rigors, and any tendency to somnolence **Meningitis** of the cerebral cortex may give rise to wild delirium, but it is less persistent than that of mania, and it alternates with a state of partial stupor The pulse is high, 130, and the temperature also—103° or more—which is not the case in delirious mania, except in very bad cases The occurrence of pain in the head, vomiting, and rigors afford further signs of differentiation from acute delirious mania

**States of Depression, or Melancholia.** Kræpelin recognises the following groups of depressive states—Simple Melancholia, Stupor, Melancholia Gravis, Paranoid Melancholia, Fantastic Melancholia, and Delirious Melancholia

**Simple Melancholia** is characterised by psychic inhibition without hallucinations and without marked delusions The patient feels depressed and his mental processes are slow He is weary, his memory is defective, he is gloomy and hopeless, and loses all interest in his pursuits and surroundings He has no actual delusions, but may complain that his life has been a failure, and that everything he has had to do with has gone wrong Sometimes there are attacks of anxiety, and there may be phobias and imperative ideas suggestive of an obsessional neurosis Suicide may be contemplated, but in this form the patient does not often actually attempt self-destruction

**Stupor.**—In this condition the psychic inhibition reaches such a degree as to produce a state of stupor The patient is apathetic, and may remain for hours in the same attitude without speaking, and taking no notice of anything Many do not take their food, but will allow themselves to be fed without offering resistance Occasionally they may utter a few detached words which indicate confused and delusional ideas These patients are all potentially suicidal,

and in spite of their apparent obliviousness of their surroundings, they are more aware of what is going on about them than might be supposed. If left unguarded for a moment they may seize the opportunity to commit suicide.

**Melancholia Gravis.** In this condition there are hallucinations and delusions. Ideas of sin play a prominent part in the delusions. The patient has led a wicked life, has cruelly treated or deceived others, or is guilty of various crimes. At other times, the distress may be centred around faults committed long ago, and apparently of a trivial character. He is overwhelmed with grief because as a child he has committed petty thefts or deceived his mother. Belief in offences against God is frequent. The patient will select some denunciatory text from the Bible and apply it to himself. He has committed the unpardonable sin, has blasphemed the Holy Ghost, and a sentence of everlasting punishment has already and irrevocably been pronounced upon him. Ideas of persecution are sometimes present. People look at him with scorn, and he is in constant danger of arrest or other punishment.

**Paranoid Melancholia.**—In this form the individual has ideas of persecution and often hallucinations of hearing voices which threaten him. He believes that his medical attendant is ruining his health, that he has, for example, been so saturated with mercury that the metal exudes through his skin, or that medicines have been administered which have destroyed his sexual powers and caused wasting of the organs concerned. He may believe that poison is being placed in his food, or that his persecutors are influencing him by means of electric currents or wireless telegraphy. Such persons may bring serious charges against their physicians or relatives, and subject them to grave annoyance if the mental condition has not been recognised. The prevailing mood is gloomy and despairing, with a strong tendency towards suicide.

**Fantastic Melancholia.**—Krapelin groups under this heading cases with a greater development of delusions and abundant hallucinations. The patient sees evil spirits, monsters, heads of animals, black men, etc. A dead friend is sitting on his pillow telling him stories, or God, Satan, or the Virgin Mary is standing by the bed. Birds and animals say insulting things to him, and his persecutors are trying to murder him. He has committed terrible sins, killed many people, and the devil has come down the chimney to take him to hell. Sometimes the delusions are of a hypochondriacal type, the patient is slowly rotting away, or is suffering from horrible diseases. He has no bowels, his heart is dead, or his brain has turned to mud. His dull despondency may be interrupted by periods of anger. At times there may be more violent states, in which the patient screams, wrings his hands, beats his head, or attacks those around him.

**Delirious Melancholia.**—Krapelin describes this as characterised by profound visionary clouding of consciousness. The patient has numerous terrifying hallucinations and confused delusions. The faces of people are distorted and strangers are taken for familiar relatives. Visions of the Virgin Mary, spirits, and devils appear. The clouds are sinking down, or the whole world is burning. Though experiencing these visions, the patients for the most part lie quietly in bed, taking no interest in their surroundings, and are scarcely capable of speaking. They pass their motions under them, and stare straight in front with vacant expression. At times they may become restless and get out of bed, wringing their hands, and manifesting great distress.

**Alternating, Periodic, or Circular Insanity** (also known as "*folie circulaire*")—This condition is less common than mania or melancholia of the intermittent type. The patients have attacks of mania or melancholia interrupted by

periods of normal mentality Cole states that the most typical form is an attack of mania followed immediately by one of melancholia and then by a normal period In other patients there may be a normal period between mania and melancholia, and in yet others mania and melancholia may alternate

### DEMENTIA PRÆCOX.

Dementia præcox is essentially a psychosis of puberty and adolescence Some writers term the disorder Schizophrenia, to indicate the "splitting" of the personality which is characteristic of the disease A hereditary influence can be traced in a considerable number of cases The onset of dementia præcox is insidious, and the condition is often diagnosed as neurasthenia or some other disorder for a long time before its true nature is recognised The following main types of dementia præcox have been recognised — (1) Simple Dementia, (2) Hebephrenia, (3) Catatonia, (4) Paranoid forms, but it should be understood that the conditions overlap and that mixed types very commonly occur

**Simple Dementia Præcox.**—The early symptoms of this form are indifference mental sluggishness, and general lessening of interest in the surroundings A school boy or girl, hitherto progressing well with studies, begins to be listless and lazy He shuns company, neglects his lessons, is unable to assimilate new facts, and shows reluctance to leave his bed Attempts to arouse him from his indifference may lead to display of irritability Associated with these symptoms are headaches insomnia, and mental depression He neglects his attire and is unclean in his person Transitory delusions occur, and there may be hallucinations of hearing voices and seeing visions A patient of the working classes fails to retain his employment for any length of time, and ultimately becomes incapable of any but the simplest manual labour, and that under supervision This state may persist for many years, death occurring from tuberculosis

**Hebephrenia.**—The onset in this form is usually more abrupt than in simple dementia præcox, though here also the patient may have suffered for some months from insomnia, headache, and listlessness The early symptoms are confusion and depression, with periods of remission, which may at first suggest manic depressive psychosis Hallucinations are common, and often these take the form of voices which abuse the patient and accuse him of immoral or criminal practices There may be visions in which the patient sees prominent persons Delusions occur which may have a paranoid colouring, the patient believing that those about him are persecuting him Sometimes they are of a fantastic character Attempts at suicide may be made Periods of depression or attacks of excitability may occur, but the prevailing attitude is that of indifference, the patient sitting idly in his chair or lying in bed apathetic and uninterested in his surroundings Maniacal outbursts may occur in which the patient shouts, screams, and is destructive Most of the cases pass on to complete dementia, but in a small proportion the course of the disease appears to be arrested, and it may be possible to employ these patients usefully under supervision Physically, there is often general mal-nutrition with impairment of digestion

**Catatonia.**—This form is usually of gradual onset, but occasionally may appear suddenly At first there is usually a mild grade of depression, and in some cases epileptiform convulsions occur during this stage The characteristic symptom is a condition of bodily immobility and muscular rigidity, which may be continuous or intermittent Attempts to move the body meet with a prompt

resistance, and produce a condition of muscular tension. The limbs may be stretched out rigidly, the fists kept tightly clenched, and contraction of the facial muscles leads to grimaces. Sometimes the general muscular rigidity is associated with a condition of stupor in which the patient remains motionless for long periods. In other cases the patient allows his limbs to be flexed and placed in any position, in which he then retains them until overcome by fatigue. Various mannerisms may be exhibited by the patient. **Echoparaxia** is a term applied to the tendency to copy any action which is performed before him, such as raising the hand. **Echolalia** is the repeating of any words or remarks which are made to him. **Negativism** is the condition in which the patient does the opposite to anything he is told to do, for instance, if told to open his hand he shuts it, and if to put out his tongue, he firmly closes his mouth. **Verbigeration**, or the senseless repetition of sounds and words in rapid succession, may be displayed, or there may be repetition of stereotyped phrases. Buckley mentions a patient who replied "30 per cent coffee" to every question he was asked. Continual laughter or weeping may occur, but many patients are mute, refuse food, and will not dress themselves. Hallucinations are often marked. Impulsive acts are characteristic. A patient may suddenly and without any warning commit some act of violence, such as assaulting an attendant or smashing furniture, and then relapse into his previous state.

**Paranoid Type.**—This form, which is characterised by the occurrence of systematised delusions, is not sharply marked off from the psychosis described under the heading **paranoia**. Hallucinations are common in which the patient hears voices accusing him of crimes. Sometimes the delusions are of an exalted type.

### ACUTE CONFUSIONAL INSANITY.

The characteristic of this state is confusion of ideas, and it differs from the manic depressive group in that emotional disturbance is not so pronounced, and physical signs are more predominant. The etiology is uncertain, but it is generally accepted that in many cases the condition is due to toxic infection or to exhaustion following prolonged mental and physical effort. Read, in an analysis of 3,000 consecutive cases of soldiers with psychoses, admitted in 1917 to Netley, found 13.3 per cent of confusional states. This was the largest group after dementia præcox, which formed 20 per cent. The patient is often badly nourished, the pulse small and of low tension, the heart sounds weak, and the temperature subnormal. Anæsthesia of the arms and legs may be present. The patient is disoriented in both time and place—*i e.*, he does not know the date, or year, and does not appreciate his surroundings. His memory is disordered, confusion of ideas is marked, and he fails to recognise objects or the identity of the persons around him. Hallucinations are present and he sees visions or hears voices. Delusions may occur, but they are not marked. At times he may become excited and incoherent.

### PARANOIA.

The characteristic feature of this disorder is the presence of *fixed*, systematised delusions. The term, however, has been used with somewhat different significations by various authorities, and there is still not complete unanimity as to what cases should be assigned to this heading. Kræpelin defines **paranoia** as the "insidious development of a permanent and unshakable delusional

system resulting from internal causes, which is accompanied by perfect preservation of clear and orderly thinking, willing and acting" <sup>1</sup> This conception of paranoia has been generally accepted, and many mental states which would formerly have been classified as paranoia are now assigned to the paranoid forms of dementia præcox or other psychoses. When the delusion is of an innocent and very restricted nature, the behaviour of the patient resembles that of a harmless eccentric individual. Clouston <sup>1</sup> mentions such a case, in which the patient was intellectually acute and morally irreproachable, but he believed that twice two were not four, but four and a quarter. He spent his whole time not devoted to keeping the asylum accounts—which he did accurately on "the old system" in deference to prevailing prejudices—in making elaborate calculations by his own system as to the distance of the stars, and in formulating new tables of logarithms, his manuscripts filled two large chests, which he solemnly left by will to the University of Oxford.

The description of the delusions of paranoia as "systematised" means that the patient rationalises his beliefs and defends his position with logical consistency. Such patients are absolutely impervious to demonstration or argument directed against their dominant ideas, and often exhibit an unmitigated contempt for the mental capacity of those who presume to differ with them. The ideas of such patients are inconsistent with the actual state of things, but the deductions drawn from these ideas are often quite rational. grant them their premiss and the rest falls in, in orderly sequence. A woman fancies that she is the Princess of Wales, she conducts herself with what she believes to be a befitting degree of dignity, decks herself with fictitious jewellery and adornments, and exacts a deferential behaviour from those who come in contact with her. A man believes that he is endowed with the power of healing by the laying on of hands, he makes his power known by inflated addresses written and oral, calling upon all who have charge of sick folk to bring them to him to be made whole.

Delusions associated with depression and ideas of persecution are very common in paranoia. Patients so afflicted often fancy themselves the victims of some occult agency directed against them by certain people, who are often not specified, but vaguely spoken of as the "villains" or "those mesmerists." One man believes that if he allows people to touch him, or even to come near him, they can read his thoughts, consequently he keeps everybody at a distance as far as he can. Another believes that there is an attempt on the part of some malicious agency to inflate him with gas, so that he will sail away like a balloon, and be lost, in consequence he will not sit down for months, but walks about, or leans against objects, ready to start away should an attempt be made. One patient believes that people are following him about, watching him, or setting traps to catch him. His letters are being opened, or his thoughts are being read. He changes his residence, but his persecutors soon find out his new address, and he recognises by various signs that their nefarious efforts to injure him are again being made. Such delusions are termed **systematised**—*i.e.*, the patient finds reasons for the attacks. In another class of cases suspicion is directed against individuals, a man suspects the chastity of his wife and believes that, with the aid of her alleged paramour, she is in some secret way depriving him of his mental and virile powers.

Those who are subject to delusional insanity may be able to conceal their delusions under the influence of a strong incentive to do so, even without any effort to disguise their ideas, if their interlocutor does not happen to touch

<sup>1</sup> *Clinical Lectures on Mental Diseases*, 1883

upon the subject of the delusion they may be able to maintain a rational conversation for a considerable time without betraying themselves. Maudsley<sup>1</sup> quotes the case of a Commissioner who was sent to an asylum to liberate those whom he judged to have recovered. He examined an old man, who gave no indication of incoherence nor insanity in any way, and an order was prepared for his release, to which he had to put his signature, he took the pen and wrote "Christ."

Hallucinations of hearing are common in delusional insanity, the patients constantly complain of voices proceeding out of the cellar and through the walls. Hallucinations of taste and smell are also frequent, and give rise to notions of being poisoned or suffocated with noxious gases.

From the medico-legal point of view, paranoia is of great importance, as the condition may result in all sorts of charges being brought against other persons, and sometimes the delusions of persecution lead to homicidal attempts upon the supposed enemies. Others put forward claims to estates or titles. Sometimes the delusions assume a religious character, and the patient believes that he has divine powers or is the Messiah. Such a one may endeavour to fulfil his supposed mission and summon meetings or distribute circulars. In certain cases the delusions have an erotic character. These are met with mostly in young women, in consequence of which they accuse men of having made improper advances to them, or worse, such ideas are sometimes very fixed, and a woman subject to them will persistently write compromising letters to the man whom she imagines has wronged her, and even to other people, making known his alleged misdoings.

Paranoia always begins insidiously, and the delusions progress slowly. The condition does not tend to shorten life, nor to pass on to weak-mindedness or dementia.

The Freudian school have found a relation between paranoia and homosexuality. Ferenczi considers that the condition is a defensive mechanism against homosexual desires in the subconscious mind, and he describes in detail several cases which support this view.<sup>2</sup>

### EPILEPTIC INSANITY.

Apart from the tendency to progressive deterioration of the intellect in epileptics, the disease may produce psychical disturbances of an acute kind, which bear a certain relation to the periodic attacks. According to Hughlings-Jackson,<sup>3</sup> epilepsy is "a sudden, rapid, excessive, occasional, and local discharge of the cerebral cortex." In the ordinary epileptic seizure, the first indication of the discharge is manifested by a sensory disturbance—the so-called aura, further recognition of development in this direction is checked by immediate loss of consciousness, the discharge then expends itself in the motor tracts. Inasmuch as the discharge may take place at any part of the cortex, the relation between the psychical and motor effect is not constant. In the ordinary epileptic seizure the psychical disturbance is limited to the aura which precedes and the lethargic sleep which succeeds it. In the cases which are of interest to the medical jurist the motor effects are negligible, and they may be entirely absent, his attention is concentrated on the disorder of the higher centres by which the mental attitude and the actions of the individual are affected. The psychical disturbances occur either before or after

<sup>1</sup> *Responsibility in Mental Disease*, 1874.

<sup>2</sup> *Contributions to Psychoanalysis*, 1916.

<sup>3</sup> *West Riding Asylum Rep.*, vol. iii.

the motor discharge, and, what is of the greatest importance from the forensic standpoint, even in its absence. During a variable period before an epileptic seizure the individual often displays a restless, irritable, morose disposition, with delusions, in which suspicion of the motives of those about him plays a prominent part, less frequently there is elation. This mental change is easily recognisable by those who are in daily contact with the patient, and from it they infer that an attack is imminent. After a fully developed seizure the patient usually sleeps profoundly for a variable period, and on awakening is in his former condition, except that he feels sore and fatigued, in some cases, automatic movements are made before consciousness returns. A man may have a fit in the street, and after it is over, may get up and walk a considerable distance without any perception of his surroundings, when he comes to himself he finds that he is in a neighbourhood quite unknown to him. Whilst in this condition he may perform a variety of actions of a purposive nature which demand complicated movements for their execution. The actions performed are determined by the influence of past co-ordinations on the motor centres acting in the absence of the restraint imposed under normal conditions by the highest controlling centres, the latter being in a state of passivity from exhaustion. Many of these actions present all the appearance of volitional movements, and yet the individual, both at the time and afterwards, is quite unconscious of having performed them, although described as automatic, they are probably initiated by a slight peripheral stimulation, or by one of internal origin. There is a tendency to resume an action interrupted by the seizure, hence a person who is walking at the moment the attack comes on, may recommence doing so as soon as the motor exhaustion is recovered from, but before the psychical centres have regained their activity.

After an ordinary epileptic seizure has occurred and the subsequent comatose sleep has passed off, the patient, instead of regaining his previous mental state, may be furiously maniacal, this condition may come on at any time within twenty-four hours or more after a fit, it may immediately succeed the coma, but more frequently some hours elapse. It may follow a single fit, but is more likely to occur after several have taken place in rapid succession. In the absence of the higher control, the lower centres are let loose and give rise to a display of blind, ungovernable frenzy which is in the highest degree dangerous to the patient and to those around, whilst under its influence the patient, without cognisance of the act, may commit deeds of brutal violence. It is important to remember that a dislike or suspicion, aroused during the stage of irritability that precedes an attack, may determine the action after it, an animus thus conceived against any one may lead to his being assaulted after the fit is over, whilst the patient is still unconscious of his actions. Occasionally the mental disturbance after an ordinary epileptic seizure takes another form, the patient being simply excitable—talking, gesticulating, and behaving extravagantly—but without violence.

The condition of "automaticity" may occur during a slight epileptic seizure without motor symptoms—the so-called *petit mal*. During an attack of this description there may be no perceptible movement of the body or limbs, the face will become pale for a moment or two and subsequently flushed, and that is all that usually is to be seen. The patient himself may or may not perceive a momentary loss of consciousness, if he is talking at the time the attack comes on, he stops and hesitates for an instant, and then resumes the conversation, if he is writing, the pen probably drops out of his hand, but the whole affair appears so insignificant that it might easily be mistaken for an

accidental slip of the fingers. If he is doing something at the time the seizure occurs, he either continues the action on the same lines, or he does something absurd or incongruous which has a certain relation to the original purpose. For example, a woman who was toasting bread had an attack of *petit mal*, she thrust the toasting-fork with the slice of bread on it between the bars and vigorously stirred the fire. This association of ideas has an important medico-legal bearing, during or after a seizure that does not produce motor disturbance there seems to be a tendency to perform some action. If at the time the person is not manually occupied, he may do something that has a strong resemblance to a purposive unlawful act, without volition or cognition, the mere sight of an object may be sufficient to suggest picking it up and putting it in the pocket. This may be done in such a way as to resemble secret theft, but the action is not unfrequently committed without attempt at concealment. Colman<sup>1</sup> gives an instance in which a man had an attack of *petit mal* whilst in an ironmonger's shop, he placed a large coal-scuttle on each arm and deliberately walked out of the shop.

A patient under the influence of an attack of *petit mal* sometimes displays another tendency which may bring him within the grasp of the law, he is seized with a desire to micturate, and regardless of the surroundings, unfastens his clothes and deliberately performs the act in public, with the result that he is taken in charge for indecently exposing his person. Colman relates the case of a woman who whilst at a public entertainment had an attack of *petit mal*, in the course of which she lifted her clothes and there and then urinated, her friends had much difficulty in convincing the police that the act was not one of wanton indecency. Occasionally the patient commences undressing in public, Gowers accounts for this on the supposition that a feeling of illness experienced after the attack "suggests" the propriety of going to bed.

A condition occasionally occurs after an epileptic seizure, called retrograde amnesia—*i.e.*, loss of recollection of all that has passed for some time *previous* to the attack. This condition may follow either an ordinary epileptic seizure, or an attack of *petit mal*, and it may occur only once out of a vast number of seizures in the same individual. Seglas<sup>2</sup> relates an instance which occurred in a young man who, whilst at dinner, had an epileptic seizure, and when he came to himself it was found that he had absolutely no recollection of the events of the morning preceding the attack. He had arranged the books in the library, had written some letters, and had paid several visits, all of which actions were completely and permanently obliterated from his memory. The patient had had many previous epileptic attacks, but had never before suffered from retrograde amnesia. The amnesic period may reach back from an hour or two, to days, or even weeks, and the conditions may be either transitory or permanent.

The cortical discharge, which under ordinary conditions determines an attack of epilepsy with clonic spasms, may apparently be entirely expended on the emotional centres, producing a state of maniacal excitement like that already described as occasionally occurring after an ordinary seizure, but without the usual fit. Individuals thus affected will commit the most brutal acts of violence, being at the time perfectly oblivious both to the injuries they inflict on others and to the damage they themselves may sustain. For two reasons, no insane condition is more dangerous than this—first, the immediately antecedent state of the individual who is subjected to its influence may be one of perfect sanity, and therefore he may be at liberty to attack the first person

<sup>1</sup> *The Lancet*, 1890

<sup>2</sup> *Annales d'Hygiène*, 1897



with whom he comes in contact without let or hindrance, second, his ungovernable fury knows no bounds, and he "goes for" his victim regardless of consequences. This condition is sometimes called **masked epilepsy**, a term which is also applied to a state of unconsciousness accompanied by automatic actions without violence.

### GENERAL PARALYSIS OF THE INSANE.

This disease is much more frequent in men than women, the subjects of it are usually strong, vigorous, and in the prime of life—from 35 to 45 years of age. Persons of a sanguine temperament are more liable to be attacked than those of a melancholic type, it is probably more common in the lower than in the higher ranks of society. A history of syphilis is present in the great majority of cases. Probably the disease is an invariable antecedent. Among the **exciting** causes are—prolonged mental strain, especially when followed by disappointment, shock to the nervous system, such as is caused by the sudden occurrence of a domestic or business trouble, less frequently brain-disturbances of a physical kind due to mechanical injury, or to sun-stroke.

The **symptoms** are divisible into **psychical** and **physical**, both kinds being highly characteristic when the disease is present in a typical form.

As in other forms of insanity, there is an incipient period which in this disease presents special features of a distinctive nature. A vigorous, healthy man, of sanguine temperament, who is fond of what is called "company"—*i.e.*, good eating and drinking—becomes more than usually jovial and loquacious. His friends probably think that he is indulging too freely in alcohol, and, indeed, the condition very closely resembles that produced by slight excess of this kind. The patient is obtrusively familiar with people with whom he has but a slight acquaintance, he will call upon them at their houses and harangue them at great length about his private affairs, he will talk by the hour about himself, his family, or his possessions in an inflated style, describing everything as the superlative of excellence. He is prone to do *outré* things, the outcome of extreme self-complacency, he will "drop in" uninvited to dine with a family with a member which he is merely on speaking terms. He becomes very forgetful, neglects appointments, and is irregular in his daily habits. An early indication, which is often the first to rouse the suspicions of the patient's friends as to his sanity, is extravagance in purchasing useless articles. If he sees a watch that pleases him, he will buy half a dozen, he will give orders to tradespeople for all sorts of goods, many times in excess of his requirements and income. Acquisitiveness may also be displayed by stupid thefts committed without attempt at concealment, the articles appropriated being often, though not invariably, useless.

The sexual proclivities usually assert themselves in a manner that demonstrates the loss of moral control, a man in the early stage of general paralysis will commit an indecent assault on a woman in a most casual way, without any of the cunning displayed in other forms of insanity. He does not appear to be driven to the act by sensual impulse, but the opportunity being present, he avails himself of it out of mere caprice without considering the consequences, just as he senselessly commits a theft. In a less objectionable way he may show his inclination towards the fair sex by proposing marriage to several women on the same day. A good deal of this folly seems to depend on impairment of the memory as well as on exaltation of the sexual feelings, though

there is an undoubted vein of lubricity present, which is evinced by a tendency to libidinous conversation

The memory and the power of concentrating the attention is always seriously impaired, even in an early stage of the disease, the mind, butterfly-like, flits from object to object, and the one on which it last settles excludes for the moment its predecessors. A good way of testing the memory is to get the patient to write a letter, if the disease is at all advanced the writer will omit words, especially short words, such as articles and prepositions

So far, although his conduct has been characterised by a variety of reprehensible actions, chiefly remarkable for their silliness, he may not have shown any decided indications of downright insanity. About this stage of the disease the **physical** symptoms are usually first observable, they may appear earlier, before the mental condition is so far advanced, or they may be delayed. The first indication is afforded by the speech, a man in the early stage of general paralysis blurs his words, or some of them, just as a person does who is slightly under the influence of drink. There is more of a hesitancy than a stammer, as though the muscles concerned in articulation co-ordinated badly, in the act of talking the lips quiver, imparting an appearance aptly compared by Bucknell to that of a person about to burst into weeping. If the tongue is protruded, fibrillar tremors of its muscles will probably be visible. The pupils are frequently unequal in size, or they may be extremely contracted or considerably dilated

As the disease progresses the mental symptoms increase in intensity, bombastic utterances are supplemented by delusions which have a like tendency. The patient fancies himself wiser, stronger, or richer than other men, his wealth amounts to millions of pounds, he has country residences in every county of England, he can outstrip the fastest horse on foot. Delusions chase one another through his mind, he recurs to the same idea, but modifies or alters its expression, and after a time he takes up other grandiloquent notions. If taxed with the absurdity of his statements, he makes little attempt to defend or justify them, but maunders on regardless of controversy, he thus differs from melancholia and other insane patients, who hold to their delusions and insist on the truth of their utterances. By this time the memory is so impaired that the patient is incapable of comparing ideas, he cannot perceive the least incongruity in his statements, they represent what to him actually exists. His face glows with transport as he recounts his heroic deeds and enumerates his vast possessions, in repose it presents the exactly opposite look of a half-demented person—stolid and fatuous. The speech is now distinctly impaired, and subsequently becomes numbing and difficult to understand

Before this stage—sometimes quite early—the patient will have suffered from **epileptiform** seizures, varying in degree from a mere “faint” (*petit mal*) up to well-marked convulsions, and even true epileptic attacks. The ribs are sometimes fractured without the patient being aware of it, due partly to trophic changes in the bones producing unnatural fragility, and partly to insensitiveness, which blunts the perception of pain

The **last** stage is one of paralysis and dementia, the patient may be able, when supported, to totter about for a little longer with a well-marked ataxic gait, and then he is confined to his bed, and lies with the knees drawn up against the abdomen, with body and mind paralysed until death occurs. The duration of the disease varies from a few months to three or, possibly, five years

**Exceptional Forms of General Paralysis.**—In the early stage, and very rarely subsequently, the condition is melancholic instead of exalted. In the ordinary

type of the disease the patient's life may be cut short soon after the physical symptoms are first developed Sankey<sup>1</sup> describes the symptoms in these cases as resembling those of acute meningitis

In a limited number of cases remissions occur, and, for a time, the patient is supposed by his friends to have recovered Blandford<sup>2</sup> states that he has seen a wonderful disappearance both of bodily and mental symptoms, the improvement lasting for some time, and that amongst these cases he has seen some which certainly would not have been pronounced insane by a jury, they had either lost their delusions, or were competent to deny and conceal them If such cases remain free from mental work, they may slowly decline without return of the acute symptoms, if they attempt to occupy themselves with their former avocations, the acute symptoms return and the relapse is progressively and quickly fatal

## TOXIC INSANITY.

### ALCOHOLIC INSANITY.

Mental disorder due to alcohol is either **acute** or **chronic**. Acute alcoholism is divisible into two types—**Delirium ebriosum**, and **Delirium tremens**.

**Delirium ebriosum**, or acute alcoholic delirium, is a condition of maniacal excitement, directly due to excess of alcohol, usually it immediately follows the bout of drinking Excess of alcohol does not necessarily imply a large amount, for various reasons, such as hereditary idiosyncrasy, previous heat-stroke or traumatic injury to the head, a greater susceptibility to the exciting effects of alcohol exists in some people than in others, therefore, the amount which is to be regarded as excessive is a variable quantity Where there is strong predisposition to become deliriously excited under the influence of alcohol, the amount taken may be relatively small, and there may be an interval of a day or more between the bout of drinking and the delirium As a rule, the acute delirium immediately follows or, rather, is a continuation and exaggeration of the wild excitement that drink not unfrequently produces in some men at the time that it is taken The condition is one of excitation of an aggressive type, with delusions of persecution or suspicion, the patient is usually clamorous, and vows that some one is trying to injure him, that his wife is unfaithful to him, and that he will kill those who have done him wrong He is usually so violent as to need restraint He has not the same kind of hallucinations as those which occur in delirium tremens—of rats and insects, but hears voices accusing him of crimes or defaming his character, or else he has sensations of a visceral type, which he says are caused by those who are persecuting him The tongue is foul and the temperature may be elevated a degree or two The mental state resembles mania more than delirium tremens Unless there is a strong predisposition to insanity the excitement passes off in a few days, though there may be hallucinations for a short time longer

**Delirium Tremens.**—This may come on in the course of a drinking bout, or some days after its cessation The patient cannot sleep, or, if he does, he dreams horrible dreams that haunt him when awake At first, he is tremulously anxious to follow his usual occupation, but cannot concentrate his attention on it, he then becomes more restless and excited, muttering and moving to and fro in a busy, purposeless manner If spoken to he lends a momentary ear

<sup>1</sup> *Lectures on Mental Disease*, 1884

<sup>2</sup> *Insanity and its Treatment*, 1884.

and answers a question, but immediately rambles off to something irrelevant, probably in the direction of his work. He stops in the midst of a muttered monologue to listen to an imagined voice, and will reply to it, he is extremely suspicious and looks and listens at all points of the compass. He constantly picks at the bedclothes, and tosses them to and fro. He sees rats, mice, or creeping things on his bed and running up the walls, which he tries to chase away or to catch with his hands. He will fix his eye on a distant spot and follow the movements of some hallucinatory creature till it reaches his bed, when he springs out, and, in a state of tremulous agitation, rushes away to escape from it. He is constantly lifting up the bedclothes, or pulling the curtains on one side to look for lurking foes. Associated with a blustering, defiant demeanour, he displays a cowardly disposition that usually renders him amenable to firm moral treatment on the part of strangers, but he is utterly intractable to attempts to quieten him made by his wife or those of his household. There is a tendency to suicide, which is often manifested by patients who have not displayed any marked degree of delirium, in the more excitable kinds of delirium tremens homicide may be committed. In the usual course the disease lasts two or three days, and terminates in several hours' sleep.

**Chronic alcoholic insanity** comes on after long-continued excess of alcoholic beverages, it is accompanied by sensory and motor affections due to peripheral neuritis, which impart a special character to this form of insanity. The mental condition is one of progressive enfeeblement, the memory fails, and the power of concentrating the attention diminishes until the patient becomes stupid and indifferent to all around him, he is negligent as regards his personal appearance, and goes about with dirty, ill-adjusted clothing. He then develops **delusions** founded on hallucinations, mostly aural—he hears people speaking evil of him and conspiring to do him bodily harm, or voices ceaselessly repeating abominable suggestions. Visual hallucinations, so characteristic of delirium tremens, are much less frequent in chronic alcoholic insanity. The hepatic and gastric troubles caused by prolonged excess of alcohol give rise to delusions that living animals are inside the stomach, or that poison is being secretly administered. The *paræthesiæ* due to peripheral neuritis are similarly misinterpreted—the patient declaring that he is being tormented by electricity administered by those who were plotting against him. It is characteristic of the change in direction taken by public thought during the latter half of the nineteenth century, that the delusions of the insane which are founded on tactual and other sensations are almost invariably referred by the sufferers to electricity—delusions which formerly have been attributed to demoniacal influence. Patients suffering from chronic alcoholic insanity are prone to delusions relating to the sexual organs, they believe that they are being subjected to treatment intended to harm them and destroy their sexual powers, either by means of poison administered in their food or by some occult application of electricity. Another frequent delusion in this relation is that the patient believes his wife is unfaithful to him, and is plotting with some one to get him out of the way.

More rarely the delusions in chronic alcoholic insanity are of an exalted type, and strongly resemble those of general paralysis. They are often equally lofty in sentiment—the patient thinking that he is possessed of boundless wealth, or that he is one of the persons of the Trinity—but they are not accompanied by the fatuous self-complacency which is characteristic of general paralysis, there is always an undercurrent of distrust as to the motives of some person or persons.

The **physical** symptoms consist of **sensory or motor** disturbances, due to

peripheral neuritis. The relation between the psychical and physical symptoms caused by alcoholic excess is not constant, an advanced stage of paralysis may exist, with but slight mental degeneration. The **sensory** disturbances take the form of extreme tenderness of the muscles of the legs chiefly, the least pressure causing acute pain, the nerve-trunks may be swollen and tender on pressure, there may also be paræthesiæ of various kinds, a burning sensation in the feet being especially troublesome. In the early stage, cramps in the calves of the legs are common. The **motor** symptoms first show themselves by difficulty in walking, the legs feeling weak, the gait, though ataxic in appearance, is really paralytic. When the patient is in bed the ankles and toes are flexed, the heels being drawn up, later on the wrists also drop. As the paralysis progresses more muscles are implicated, including those necessary for the carrying on of respiration, and death results.

The drinking of alcohol in excess is not only a cause of insanity, but it may also be the result, the term **dipsomania** has been used to signify a morbid tendency to take drink. The habit of drinking to excess is but too easily acquired by those who have no special proclivity to it, but there are undoubtedly cases in which it is directly due to hereditary transmission, it may also be the result of the emotional state and enfeeblement or moral control which marks the onset of many forms of mental disease. In addition to the immediate effects of alcohol on the mind, a further and debasing change is produced in the moral character, the individual loses both self-control and all regard for truthfulness, he will resort to any subterfuge to gratify his cravings, and will lie in a most barefaced manner to extricate himself when in a difficulty. Such people are pests to their families and to society at large.

The habitual use of **morphine** produces a still more debasing effect on the moral character. A dipsomaniac may pull himself together for a short time and abjure drink, but rarely does an opium *habitué* voluntarily cease taking the drug, his craving for it is such that nothing short of physical hindrance will restrain him. Under the sway of this habit, men, and even women of refined character, will resort to the meanest artifices to gratify it. If the habit is of long continuance the memory becomes impaired, this at first seems to be the result of inattention, the patient is absent-minded and does not give himself the trouble of focusing his thoughts upon the subjects under consideration. At this period, if sufficient mental stimulus is brought to bear, he is capable of concealing his weakness by increased mental effort, subsequently, there is an actual loss of memory which cannot be overcome. In this stage, hallucinations are common and may give rise to delusions like those of chronic alcoholic insanity. The physical indications are loss of flesh, tremulousness when not under the influence of the drug, anorexia, or capricious appetite and diminution in muscular power. A very significant symptom consists in recurrent visceral neuralgia, which may resemble the agony caused by the passage of a gall-stone, gastric crises (mistaken for bilious "attacks") are common. The habit is exceedingly difficult to eradicate, but, if eradicated, recovery is more complete than in the analogous condition produced by chronic alcoholism, because of the absence of vascular and other degenerative changes which accompany the latter, and which, of course, are permanent. It is not unfrequent for those who have been cured of the opium habit to take to drinking. If mental enfeeblement through opium-eating is pleaded as a bar to criminal responsibility it is to be regarded much in the same light as the plea of chronic alcoholism.

**Cocaine**, when taken habitually and for a long time, produces disastrous effects on the mental powers. The evil is usually initiated by the use of the drug

as a therapeutic agent for the relief of pain, or irritation. Ere long the habit is acquired, and the victim becomes apathetic and feeble both in body and mind, although directly after recourse to the drug he will be complaisant and even bright for a time. His sleep is fitful and troubled, and his appetite forsakes him. He becomes profoundly melancholic, and tends to develop delusions of persecution, and to be disposed to commit suicide. Like morphine, cocaine eradicates the moral sense, and degrades its victim into a shameless prevaricator.

Chronic **lead-poisoning** may give rise to certain psychoses. The symptoms produced do not appear to be always the same. Savage<sup>1</sup> observed acute mania in one case, and in another, symptoms closely resembling those of general paralysis. In a case seen by the author there were delusions with depression.

**Psychoses associated with Somatic Disease.**—Many bodily diseases may be accompanied by mental disturbance. Acute delirium may be present in the exanthemata, and other acute conditions associated with high temperature. In states of exhaustion a low muttering delirium is frequently present before death. Delirium may arise during the acute stage of influenza, and after the bodily symptoms have subsided there may be a state of depression which may pass into melancholia, or sometimes confusional insanity with excitement and hallucinations. Chorea is often characterised by mild symptoms such as dullness, inattention, and loss of memory. In some cases attacks of acute confusional insanity occur. In myxœdema the patient is often dull and lethargic, and sometimes there are attacks of confusion with hallucinations and excitement, which may be followed by a state of stupor. In hyperthyroidism, restlessness is common, and there may be attacks of excitement, with insomnia, hallucinations, refusal of food, and violence. In uræmia there may be mental wandering, with hallucinations and agitation. Syphilis, in addition to general paralysis of the insane, which is described separately, may give rise to confusional insanity or dementia, following involvement of the membranes or vessels of the brain. Other diseases in which mental symptoms may be present are arterio-sclerosis, diabetes, advanced disease of the heart or lungs, and paralysis agitans. Injury to the brain does not produce any definite form of mental disorder, and the differentiation of traumatic insanity is not justified. Mental disturbance following a blow on the head is usually associated with concussion, and most cases clear up completely. Sometimes, however, there are permanent changes, and the individual suffers from irritability, confusion, and lapse of memory. There is reason to believe that trauma may accelerate the onset of other forms of insanity, such as G.P.I. or mania, but its most serious effects are probably experienced by alcoholic subjects in whom alcoholic insanity may be precipitated.

### SENILE DEMENTIA.

Senile dementia is not an invariable accompaniment of advanced age, and therefore no limit can be assigned at which the mind begins to give way from physiological decay. Some men retain their mental vigour in a high degree until far advanced in life, they display a wonderful power of abstract reasoning, and of memory for words of highly specialised meaning, such as proper names, which being least organised, soonest undergo dissolution. A man of less mental vigour shows signs of decay and loses coherence of ideas ten or twenty years sooner. If signs of dementia occur before the age of sixty years, there has probably been some cause other than age which has interfered with the nutrition

<sup>1</sup> *Insanity and Allied Neuroses*, 1884.

of the brain, defective memory frequently betrays itself at a much earlier period, but the reasoning powers remain active. A certain dulness of intelligence, as compared with that which formerly existed, is frequently the precursor of dementia. In this stage, a man of considerable attainments will be unable to solve at the moment a question involving abstract thought, which would formerly have given him little trouble, if the question is written down, and he is so minded, he will arrive at a correct answer when left to himself for a time, the power of abstract reasoning is dulled, but not destroyed. This applies, though not so directly, to minds of lesser culture, and should be remembered when examining cases which are supposed to display great mental weakness.

When dementia follows a progressive course, the mental boundaries may slowly contract without other change than gradual intellectual dissolution. In some cases, phases of excitement or of depression occur at intervals, there is always more or less loss of self-control, and frequently excessive irritability of temper. Hallucinations and delusions are not uncommon, and there may be a tendency to suicide. Erotic impulses may be present, which are not unfrequently the cause of unchaste behaviour on the part of old men to young girls.

Dementia may be the final stage of any of the forms of ordinary insanity, such as mania or melancholia, this is known as **secondary dementia**. The condition is that of mental enfeeblement, advancing to one of complete abolition of intelligence. There may be occasional waves indicative of the initial psychosis, the patient being excited or depressed at intervals. The animal propensities, uncontrolled by reason, may display themselves, or the patient may be impassively tranquil unless annoyed, when he blazes up into a momentary fit of passion.

### INSANITY ASSOCIATED WITH REPRODUCTION.

There is no definite form of insanity associated with the puerperal state, but under the strain of pregnancy or parturition a woman predisposed towards mental disorder is more likely to experience an attack of manic depressive insanity or some other psychosis. There are, however, often special features displayed in the so-called condition of puerperal insanity, and it will be convenient to consider these as they are manifested during pregnancy, puerperism, and period of lactation.

**Insanity during Pregnancy.**—Emotional disturbance is a well-recognised accompaniment of the period of gestation, it is manifested by strongly marked likes and dislikes in relation to food, and not unfrequently by alteration in the general mental state. Some women are morose or irritable, others are usually complaisant during the period of gestation. If there is any neurotic taint, or even in the absence of discoverable hereditary proclivity, such emotional disturbance may exceed the limit recognised as compatible with self-control and may develop into actual insanity. Insanity during gestation is exceptional, when it occurs, it is usually after the third month. The condition is one of depression, accompanied with suspicion and dislike of husband and (should the patient be a multipara) of children, this may alternate with states of maniacal excitement, or the entire disorder may be of the exalted type. Mental disorder is most liable to occur in first pregnancies. There is always a strong tendency to **suicide**, and there may be to **homicide**. The patient sleeps little, and is subject to hallucinations and delusions.

**Insanity during the puerperium**, when it occurs, usually comes on within a fortnight after parturition, in its most frequent phase it partakes of the **exalted type**. The patient is sleepless for a night or two, and then is excitable and talks more than usual. She is capricious and, without any reason, takes a dislike to those about her, she gives the nurse all the trouble she can by demanding a number of trifling services and then complains of her inefficiency. She displays the greatest antipathy towards her husband and her child, and declines to speak to the doctor, or roundly abuses him to his face and calls him by some opprobrious epithet. Food is obstinately refused, necessitating forcible feeding. The delirium, accompanied by hallucinations and delusions, is very acute, and is often characterised by obscenity of language and gestures. The mental state may be one of **depression** in place of exaltation, the patient being profoundly melancholic, in this state she takes no notice of anything, perhaps not even speaking for days together. In both forms the **suicidal tendency** is strongly developed, and as the patient is impelled to the act by overpowering impulse, originated by vivid delusions, there is always the probability of a sudden outburst, even during a period of temporary quiescence. The same observation applies to the **homicidal tendency**, which is also common in puerperal insanity, and is chiefly directed against the child or children of the patient, the temptation to take her child's life is so strong, that, although the woman's instinct rebels against it, she feels herself incapable of resisting, and will therefore sometimes ask that the child may be kept away from her.

**Insanity during lactation** may occur from the third to the sixth or eighth month after parturition, or even later. It is essentially due to enervation with, probably, some substratum of neurotic strain, if there is any predisposition to mental derangement, the sequence of gestation, parturition, and lactation causes it to develop. The insanity of lactation usually commences by loss of sleep, depression of spirits, delusional hallucinations, restlessness, and fretfulness. These symptoms may deepen into melancholia with or without outbursts of excitement, or a maniacal condition may predominate. Bevan Lewis<sup>1</sup> states that the more acute forms of excitement prevail within the first three months following parturition, and delusions of persecution, with the associated gloom and despondency of melancholia, when the mental symptoms first betray themselves, six or more months after. The impulse to **suicide** and **homicide** is very great.

### WEAK-MINDEDNESS.

Mental weakness may be developmental, or sequential. Idiocy, imbecility, and senile dementia belong to the first group, and dementia as a terminal stage of any of the forms of simple insanity, or when caused by coarse brain-lesions, to the second.

### IDIOCY.

Idiocy implies a condition in which there is either congenital deficiency of mind, or a deficiency caused by processes which take place after birth, but before the period when the mental faculties begin to develop. **Imbecility** is the same condition in a lesser degree. Lawyers include both forms under the term *dementia naturalis*. Idiots may be roughly divided into micro-cephalic and megalo-cephalic.

<sup>1</sup> *A Text-Book of Mental Diseases*, 1889



The **micro-cephalic**, or small headed, idiot is of the congenital type Ireland<sup>1</sup> states that the size of the head affords no evidence of the comparative intelligence of the idiotic child, but if the head is less than 17 inches in circumference the intellectual power will be feeble. The **megalo-cephalic**, or large headed, idiot belongs, for the most part, to the group which results from pathological changes after birth. Rickets (?), syphilis, and hydrocephalus are the diseases chiefly credited with the causation of this form of idiocy. Hypertrophy of the brain, and more especially hydrocephalus, may be congenital. Bury<sup>2</sup> shows that syphilis may lead to hindrance of brain growth from thickening of the cranial bones caused by osteitis in early life, from thickening of the membranes and the coats of the arteries, and from sclerosis of the cortex, all of which usually come into play after birth, congenital deficiency of mind from inherited syphilis is comparatively rare. Idiocy may be initiated by convulsions, which often occur at the time of teething. Traumatic injury, such as the head sometimes sustains during difficult parturition, may give rise to imbecility or idiocy in combination with spastic paraplegia, this is due to meningeal hæmorrhage causing compression of parts of the cortex, which become flattened and atrophied. Idiocy may be accompanied by epilepsy. In the ordinary idiot the palate is narrow and highly vaulted, there are rarely more than twenty eight teeth. Pronounced bodily deformity—**cretinism** and **myxœdema**—is sometimes associated with mental deficiency—the former with idiocy, and the latter more frequently with imbecility.

Idiocy comprises all grades of mental deficiency, from a state of mere automatic existence, with considerably less intelligence than that displayed by domestic animals, upwards, through the lesser degrees of imbecility, to the confines of normal mental development. The mental condition of many persons classed amongst the sane is scarcely, if at all, distinguishable from the milder forms of imbecility.

Idiots of the lowest type are absolutely devoid of any intelligence, they are incapable of attending to their natural requirements, and they pass their time in rocking themselves to and fro, or moving their limbs in a purposeless manner. If slightly less debased, they display a marked tendency to cruelty—they will torture insects and small animals in a diabolical manner. They are mischievous, dirty, and utterly without sense of shame, some are irritable and dangerous to those whom they have power to harm. Imbeciles of a less pronounced type often give rise to an infinity of trouble by running away from home and associating with the criminal classes. As they grow up they show depraved tastes, they drink, gamble, are riotous, and are addicted to the company of prostitutes and thieves, along with this shameless depravity there is a considerable amount of low cunning and of knowledge of a certain type. Individuals of this class, or a little above it, who display a degree of sanity, often come under the notice of the medical jurist in relation to their fitness to manage their own property and affairs. A totally different type of imbecility is characterised by a gentle tranquillity of conduct, the mental defect showing itself in a purely negative manner.

### “MORAL” INSANITY.

Alienists no longer include the condition formerly described as moral insanity under the psychoses. Patients of this class are now regarded as moral imbeciles, and form Class (d) under the Mental Deficiency Act, 1913—*i.e.*, “persons who from an early age display some permanent mental defect coupled with strong, vicious, or criminal propensities, on which punishment has had little or no deterrent effect” (see p. 324). Tredgold<sup>3</sup> gives the following description of this type—“Physically, moral imbeciles are usually well-developed and present none of the signs so commonly seen in ordinary aments. Most of them

<sup>1</sup> *Idiocy and Imbecility*

<sup>2</sup> *Brain*, 1883

<sup>3</sup> *Transactions of the Royal Society of Medicine*, 1921

are quite pleasant-looking persons, indeed not a few are unusually engaging and attractive. Similarly with regard to their intellectual attainments and ordinary intelligence, moral imbeciles present no distinguishing features. They will converse well and even brilliantly upon the ordinary topics of the day, whilst many of them are so plausible in their speech and so apt at repartee as to appear exceptionally clever and astute." Nevertheless, moral imbeciles will thieve, lie, cheat, and commit sexual offences and assaults without shame or remorse. They are selfish, lacking in affection, and callously disregard the rights and feelings of others. Tredgold points out, however, that with regard to their crimes it is worthy of note that, in spite of their cunning in making elaborate plans, they will often neglect some simple precaution which could hardly be omitted by a person of average common sense, and this omission leads to their speedy detection.

In moral imbecility the higher levels of cerebral development are either imperfectly evolved from birth, or after evolution become diseased and more or less functionless, the result being that, although the intellectual capacity is not seriously affected, the emotional and automatic functions have more than normal play. Such constitutes the mental condition which characterises certain criminals, and here comes the difficulty—viz., to distinguish between moral depravity and moral imbecility, for the former merges into the latter without any absolute line of demarcation. The question has been discussed from extreme standpoints—one is, that the mere commission of a flagrant crime without remorse demonstrates moral insanity, the other is, that unless the reasoning faculties are so far weakened that the person who commits the crime cannot distinguish between right and wrong, he is responsible for the act, in other words, he is not insane. To countenance the first view is to place a premium on depravity. That individuals whose moral control is so far weakened as to place them in the ranks of the insane commit crimes of the vilest nature without remorse is perfectly true, but to accept the crime as a proof of insanity is another matter. The second opinion, which is held by lawyers, will be discussed presently.

Without entering into a discussion as to how far the emotions may determine the actions without implication of the reasoning powers, experience necessitates the conclusion that for all practical purposes, the moral perceptions may be so far perverted as to render the individual irresponsible for his actions without signs of mental disease being present, either in the form of delusions or of intellectual enfeeblement. By this it is not meant that the intellect is absolutely intact, but that it is not weakened so far as to betray its condition, except by being unable to control the moral sense.

The following instances of "moral insanity" described by the author are retained, though it is doubtful whether the second should be regarded as a typical case.

A short time ago the author was consulted with regard to a young lady of only fourteen years of age, who had been "withdrawn" from a boarding school for repeated thefts from her fellow pupils, although she was abundantly supplied with pocket-money. She had not the least regard for truth, and took a malicious pleasure in false statements which brought trouble upon innocent persons. When at home she behaved indecently with her own brother, and made improper overtures to her father's groom. Intellectually she was bright and clever, her schoolmistress reported that she was able to keep pace in classes with other girls of her age, but was fickle and lacked application. The evil strain came from her mother, who took to drinking after she was married and eventually eloped with a man of low social position; the girl was born about a year after the mother gave way to intemperance. This is an instance of defective evolution of the higher mental levels;

the finer traits of character were entirely absent, but there was no lack of a certain cleverness, which conveyed the impression that if the girl only had application and perseverance she could acquire as much knowledge as any of her compeers, moral lapses apart, her friends regarded her as sane enough

Hack Tuke<sup>1</sup> relates a very typical case of moral insanity, of which the following is a condensed abstract —A man, who in his youth was sullen, uncommunicative, idle, sly, and treacherous, at an early age evinced a disposition to torture domestic animals and to treat cruelly younger members of the family. On one occasion he took a younger brother into the fields, undressed him, beat him with long lithe willows, and bit and scratched him about the arms and upper part of the body, threatening to kill him with a table knife if he cried out. Shortly after he was apprehended for cutting the throat of a horse belonging to a neighbour, and confessed that he had maimed several other animals, and had twisted the necks of fowls and then concealed them in wood piles, he was sentenced to twelve months' imprisonment. On his discharge from prison he attempted to suffocate a little child by piling clothing on the top of it, he then stole some money from his father's desk, for which act he was sentenced to seven years in a penitentiary. After his liberation, being again at home, he saw his father accidentally cut his hand so that it bled profusely, this seemed to excite him, and he went to a neighbouring farmyard and cut the throat of a horse, killing it. He escaped, and, whilst hiding in a wood, saw a young girl, seized her and committed a criminal assault on her. After being about ten years in prison for this offence he was set free, and on his way home from prison he caught a horse, tied it to a telegraphic pole and mutilated it in a shocking manner, cutting a terrible gash in the neck, another in the abdomen, and taking a piece off the end of its tongue. For this he was tried and acquitted on the ground of insanity, and was transferred to an asylum. After being there for five years he made his escape, and was only absent from the asylum about an hour, when he overtook and attempted to outrage a young girl almost in sight of the pursuing attendants. Beside all this he was guilty of innumerable acts of cruelty to fellow patients in the asylum, and also to dogs, cats, and fowls, he was a great coward, and was never known to attack any person that would be likely to offer resistance. The sight of blood had a strange effect on this man, his face grew pallid, he became nervous and restless, he lost control over himself, and unless watched indulged in the proclivities for which he was notorious. If so situated that he could not indulge his evil propensities he was a quiet and useful man, he had had a fair education, and enjoyed reading the newspapers and letters sent to him.

### "IMPULSIVE" INSANITY.

Alienists do not now recognise a definite form of insanity characterised by the performance of impulsive acts, though the term "impulsive insanity" is still occasionally used by lawyers in criminal cases. Sudden impulses may arise and be acted upon in manic depressive insanity and other forms which have been described in the preceding pages. The following paragraphs, which were written by the author at the time when impulsive insanity was separated from other abnormal mental states, have been left because they contain many observations of value, even though the cases would now be classified under other headings.

**Homicidal Impulse.**—A woman develops an impulse to kill all her children, and resists it for a time, possibly thinking about the matter for days, that is to say, the thought keeps continually recurring. At last she acts, but, after killing one of her children, control is re-established, she throws away the knife and bursts into tears. Such a case was tried before Lord Blackburn<sup>2</sup>, the woman, after cutting the throat of one of her children, was checked in her homicidal career by the child next in turn to be sacrificed throwing her arms round her mother's neck and speaking to her in a caressing manner. The woman subsequently explained that she was going to kill the child, but that, after the caress, "she had not the heart to do it." Although she evidently knew right from wrong, and also the character of the act, the judge told the jury that there were exceptional cases, on the strength of which ruling, a verdict

<sup>1</sup> *Journ. of Mental Science*, 1886

<sup>2</sup> Cited by Orange, *Journ. of Ment. Science*, 1884

of not guilty, on the ground of insanity, was returned Maudsley<sup>1</sup> relates a very instructive example of homicidal impulse

A medical practitioner was admitted to an asylum on certificates which stated that he had made a murderous attack on his mother in law, whom he usually respected and loved. He had an attack of mental disorder when he was twenty two years of age, a second after an interval of fourteen years, and a third (the one for which he was now placed under restraint) after a further interval of four years and a half. Between the attacks he successfully conducted a large medical practice, and was so much respected by his fellow-townsmen as to be elected to the office of mayor. During his second attack of derangement he shot a gentleman with whom he was out shooting, and, although the coroner's inquest resulted in a verdict of accidental death, there were some who thought differently. He remained in the asylum for four months, during the whole of that time he betrayed no symptom of mental disease, and consequently was discharged as recovered. Twenty days after leaving the asylum he killed a female servant by cutting her throat with a razor, having shown no indication of insanity up to within a few hours of the act. Acquitted at his trial on the ground of insanity, he was sent to Bethlem Hospital as a criminal lunatic. The medical officer, after an observation of several months, said he could not attach any particular symptom of insanity to him, and that, supposing he was a private patient and the Commissioners in Lunacy asked why he was detained, he could give no definite reason for his detention.

In another class of cases the homicidal impulse exists along with, or rather in consequence of, pronounced delusions. A man falsely believes that a certain person has done him some great wrong, and, in consequence of this delusion, he murders him. Such cases are a common result of paranoia, and are usually easy of recognition, as the insane person makes no secret of his delusion. Occasionally, however, an extraordinary capacity is displayed to conceal delusions, which are yet sufficiently powerful as eventually to lead to a homicidal outbreak. Maudsley thinks it probable that the doctor previously described was able successfully to conceal his delusions when he had a strong motive to do so.

**Suicidal Impulse.**—According to the popular notion, the suicidal act in itself constitutes evidence of insanity. This is fostered by the stereotyped addendum to the verdict of the coroner's jury of "temporary insanity," probably devised as a means of escape from the verdict of *felo-de-se*, which formerly carried with it certain penalties. The suicidal act is not necessarily indicative of insanity, it may be the result of a rational resolve. If a man of high social position, and, by repute, of great wealth, is discovered to have embezzled money or forged monetary documents, and is stared in the face by the inevitable consequences, he may prefer immediate death to penal servitude. The misery—mental and physical—he seeks to avoid is real and tangible, the question is one of balance of evil, the remedy is a desperate one, it is true, but so is his condition, and the outcome is, that he determines to avail himself of the only means of escape at his command, and puts an end to his life. Probably not more than 30 per cent of suicides are actually insane, so far as any trustworthy evidence goes.

Like homicidal impulse, insane suicidal impulse may exist either with or without delusions or other disorder of the mind, the impulse may be controlled for a time, or the patient may ask to be protected against himself. Suicide from impulse differs from that with delusions, inasmuch as the patient appears to be sane on all but the one point. The tendency may be recurrent, the patient being free from mental disorder meanwhile. Instances have occurred in which an unsuccessful attempt to carry out the intention has been sufficient to restore control, and free the patient from the impulse. A case is related of a man going

<sup>1</sup> *Responsibility in Mental Disease.*

towards a river with the set purpose of drowning himself, he was attacked by thieves and defended himself lustily, and, having got away, he returned home cured of his suicidal impulse. The impulse may take a certain definite direction to the exclusion of all others, a man will neglect knives and razors, or other means at hand, to go a considerable distance and place himself on a line of rails and allow a train to pass over him. The impulse once fully developed may recur, even after a long interval of apparently perfect sanity. A case is recorded in the Commissioners' Blue Book (quoted by Newington<sup>1</sup>) of a man who had been fourteen years under supervision, for the first part of that time he had been absolutely suicidal, but he improved, and used to go out and enjoy himself. At the end of fourteen years his mother sent him an old writing desk, in it was a secret drawer, from which he took a bottle containing poison that he had placed there fourteen years before, he did not swallow the poison at once, but walked off into a wood, and there took it and died. Before the arrival of the desk, he had plenty of opportunities of procuring poison had he been so disposed.

When the impulse to suicide is the outcome of melancholia or other mental disorder, there is usually a definite delusion, the patient hears a constantly recurring command bidding him to kill himself, or accusations of horrible crimes are ceaselessly made by voices from the sound of which he seeks, in death, to escape. The delusion of persecution is a common cause of impulse to suicide. Heredity exercises a powerful influence in predisposing to suicide, the tendency often asserting itself about the same time of life in the child as in the parent. Ill-health reduces the higher inhibitive powers, and if there is a latent proneness to suicide, it is then likely to assert itself, the same remarks apply to business worries, loss of money, and disappointments of all kinds. In the latter instances the impulse is of a mixed origin, and approaches in character that which determines the suicidal act in a sane person. If a man labours under the delusion that he is the victim of a secret society, whose agents are persecuting him and rendering his life unbearable, and, in consequence of this delusion, he puts an end to his life, the act is that of an obviously insane person, because he believes a delusion and acts upon it as though it was a real occurrence. If a man has lost a great deal of money and is consequently obliged to retrench his household expenses, he may take such an exaggerated view of his difficulties as to develop an impulse to suicide, although a delusion determines the impulse it is one that is founded on an actual occurrence, and would not have come into play under more favourable external relations. The man with hallucinatory delusions is liable to develop the suicidal impulse in spite of his surroundings the other only in consequence of them.

The approach of the wave of depression which allows the impulse to develop may be felt by the patient, in some instances he requests that he may be taken care of until the depression is passed. Further evidence of consciousness of loss of self-control is afforded by the fact that, in some instances, the patient himself will lock up knives and other instruments, which from propinquity might tempt him, this teaches an important lesson on the subject of irresponsibility. Suggestion may exercise a powerful influence on a predisposed mind, hearing of the suicide of an intimate friend, or even of that of an individual personally unknown to the patient, may be sufficient to incite him to the same act. The impulse may be controllable at one time and not at another, which further complicates the question of responsibility. Threats to commit suicide made by a person suffering from some distressing bodily ailment are not unlikely

<sup>1</sup> *Journal of Mental Science*, 1886

to be put into execution Manning<sup>1</sup> relates the case of a perfectly sane man who was admitted into hospital for liver disease and dropsy. He told his wife that, if he ascertained from the doctor that his disease was incurable, he would commit suicide, and that he had at first thought of shooting himself with a pistol, but, considering that it would be a cruel shock to his fellow-patients in the ward, he had resolved to adopt another method. One day after this conversation, he learnt on enquiry from one of the resident medical officers that his case was hopeless, he watched his opportunity, went on to a balcony outside the ward, and deliberately flung himself over and broke his neck.

**Medico-legal Relations of Suicide.**--The law regards the suicidal act as felonious, unless the individual is held to be irresponsible on the ground of insanity. A person who aids and abets another person to commit suicide is guilty of murder, the crime is not lessened if two persons mutually agree to commit suicide at the same time and in the presence of each other. If one dies and the other survives, the survivor is guilty of murder, the fact of his making a simultaneous attempt on his own life does not affect the crime committed against the deceased. If a person, in the attempt to commit suicide, occasions the death of another person (he himself recovering) he is guilty of manslaughter. A man jumped into a canal with the intention of committing suicide, but was rescued by a passer-by, who unfortunately was drowned in the act of rescue, the intended suicide was found guilty of manslaughter in spite of an attempt to prove him insane (*Reg v Gethercole*).

**Kleptomania**, or impulse to steal, is of dubious existence apart from an insane condition due to general mental disease. That an insane person may have a raven-like propensity to steal is well-known, but that, because a person steals without a transparently obvious motive for doing so, he is, therefore, to be regarded as insane is simply to encourage vice. If the propensity to steal is really due to an insane state, there will be other evidences of mental disease apart from the act of theft, a person will not be absolutely sane in every respect save the desire to appropriate articles that do not belong to him. Acts of theft, we have seen, are not uncommon in the early stage of general paralysis, or after an attack of minor epilepsy, and they are usually traceable to their causation. Moral insanity with impulse, which is usually alleged in defence of the culprit when kleptomania is brought in question, is to be regarded with great suspicion. It is, of course, quite possible that a person who is the subject of moral insanity may pilfer, as well as do other acts contrary to the moral code, but it is improbable that loss of the highest control will solely exhibit itself in such a narrow channel—there will be other evidences of its existence. When kleptomania is pleaded in defence, it is usually manifest that no evidence worthy of the name is advanced to prove antecedent disorder of the mind, the most that is urged is that the accused was quite capable of paying for the stolen articles, and, therefore, that the act was due to an insane impulse.

<sup>1</sup> *Journal of Mental Science*, 1886

## CHAPTER XXVII

**MEDICO-LEGAL RELATIONS OF INSANITY.**

THE subject of insanity comes under the notice of the medical jurist in relation to (a) **criminal responsibility**, (b) **lunacy certificates**, (c) **commissions of enquiry**, (d) **testamentary capacity**, (e) **feigned insanity**.

**(a) CRIMINAL RESPONSIBILITY.**

The commission of a criminal act involves punishment, but if the person who commits a criminal act is proved to be insane at the time he committed it, he is held to be irresponsible to the law for what he has done. The questions that arise out of this statement are — What constitutes insanity, and to what extent must the accused be under its influence in order that he should be held irresponsible?

No definition of insanity has ever been formulated that can be applied as a touchstone to the individual, and his sanity or insanity thereby determined. In order to meet the difficulty, certain features assumed to indicate insanity have been selected by the administrators of the law and converted into a test, to be applied to each individual case. This "**legal test**" was enunciated in 1843 by fourteen judges in answer to certain questions put to them by the House of Lords, the occasion being the acquittal of M'Naughton on the ground of insanity, for the murder of Mr Drummond (see p 307). It is sometimes termed the "**M'Naughton decision**," and is as follows —

"To establish a defence on the ground of insanity, it must be clearly proved that at the time of committing the act the party accused was labouring under such a defect of reason from disease of the mind as not to know the nature and quality of the act he was doing, or, if he did know it, that he did not know he was doing what was wrong."

The first points that present themselves in relation to this test are — That mental oblivion as regards certain external relations is assumed to be the necessary accompaniment of an exculpatory degree of insanity of all kinds, and that its occurrence alone affords evidence of irresponsibility.

A slight acquaintance with the various forms of insanity is sufficient to show the inadequacy of the terms laid down to determine irresponsibility. A man may be so dominated by an impulse to homicide as to carry it into effect in spite of the knowledge that he is doing wrong and, therefore, is liable to punishment. At the Shropshire winter assizes in 1885 a man called Ware, who had been a patient in an asylum, and had killed another patient with an iron bar, was indicted for murder. It was proved by the depositions that the prisoner was perfectly aware of the nature of the act he had committed at the time he committed it, he acknowledged that he had killed some one, and demanded a promise from the attendant that he should not be punished if he gave up the iron bar. The case did not come before the jury, as the man was previously removed to the criminal lunatic asylum at Broadmoor, but Mr Justice Hawkins made some remarks which are very pertinent to the

question at issue - ' It would be impossible to say that Ware did not know that he had killed a man, because he said himself that he had, and it would be impossible for anybody to urge that he did not know it was wrong, for he wanted a promise that he should not be punished, but unless one put a totally different construction on the law, *that* would have to be proved, although no man in his senses would suppose that any jury would find Ware responsible for what he had done '

It is well known to medical men that a man may be under the influence of a delusion or of an insane impulse which urges him to commit an act that he knows is punishable by law, if the delusion or impulse is sufficiently powerful to overcome his self-control he commits the act—that is to say, he is unable to restrain himself. Loss of self-control from mental disease is equally causative of irresponsibility as is want of knowledge between right and wrong, therefore any one subject to such loss of control ought not to be held answerable for his actions. The legal test as interpreted by almost all the judges will not admit of this. In *Reg v Cole* (C C C, 1883), for murder of a child, Denman, J., allowed " that it was established in evidence that the prisoner had been suffering from delusions (that there were men under the floor, and in a cupboard, who sought to injure him), but that he knew that he was doing wrong, and he knew that he acted contrary to the law of this country "

In his *History of the Criminal Law of England*, 1883, Sir James F Stephen puts a much more liberal interpretation on the legal definition of insanity as a plea for criminal irresponsibility - In his opinion the law allows that a man, who by reason of mental disease is prevented from controlling his conduct, is not responsible for what he does, and that the existence of any delusion, impulse, or other state which is commonly produced by madness, is a fact relevant to the question whether or not he can control his conduct. The late Lord Chief-Justice Cockburn did not view the law in this light, but he expressed himself to a select committee of the House of Commons to the following effect—" I have always been strongly of opinion that, as the pathology of insanity abundantly establishes, there are forms of mental disease in which, though the patient is quite aware that he is about to do wrong, the will becomes overpowered by the force of irresistible impulse, the power of self-control when destroyed or suspended by mental disease becomes, I think, an essential element of (ir)responsibility " If the law thus stood, as one judge maintains it does, and another states that it ought to do, the sentences of the higher criminal courts would not subsequently be annulled, as they often are, by the influence of public opinion

The **medical view** is, that a man who is the victim of mental disease may know that a certain act is wrong, and is punishable by law, but that an insane impulse, whether arising from a delusion or not, may overcome his self-control and he may commit the act, not because he does not know that he is thereby doing wrong, but because he cannot prevent himself from doing it

**Procedure in the Case of Criminal Lunatics.**—If a man who commits murder is arrested, he is taken before the magistrates, and no matter how insane he may be he is committed for trial at the assizes. If the mental disorder is of such a decided nature as to render it unadvisable that he should remain in jail, he may, by order of the Secretary of State, be removed to an asylum, such cases, of course, could give rise to no difference of opinion at the trial. The cases which admit of difference of opinion are not sufficiently pronounced as to be removed, and the prisoner probably remains until the trial without anything being said, or any steps taken, in relation to his mental condition. It is to be



noted, however, that when, after committal, insanity is alleged, the public prosecutor if informed will direct an investigation as to the state of the prisoner's mind to be made by two medical men of experience, with whom an official referee may be associated. It does not follow that at the trial these gentlemen will be retained on behalf of the Crown, although the Home Office orders and pays for the investigation, in some instances, one or more of the examiners have been subpoenaed for the defence, their evidence being favourable to it. Even if, under the authority of the public prosecutor, the prisoner has been examined as to his mental condition, some judges will only allow the medical witness to state what he saw and heard during his interview with the prisoner, they will not admit any expert evidence, ruling that the jury alone is competent to express as to his sanity or insanity. Other judges accept expert evidence.

When the case reaches the assizes the accused is taken before the grand jury, whose function is to find a true bill, or not, irrespective of the condition of the prisoner's mind. The grand jury has nothing to do with the question of sanity or insanity. If a true bill is found the accused is arraigned before the judge and the petty jury and is called upon to plead, should his mental condition make it doubtful as to whether he is in a fit state to plead, the jury, from evidence brought before it, finds that he is, or is not, capable of taking his trial. If the verdict of the jury is that the accused is insane and is not in a fit state to plead, he is forthwith ordered by the court to be detained during His Majesty's pleasure, on the other hand, if the jury is of opinion that the accused is capable of pleading he is tried in the usual way, but this verdict that the accused is capable of pleading does not prevent another jury from finding that he was insane when he committed the deed. The Trial of Lunatics Act, 46 & 47 Vict. ch. 38, provides that if it appears to the jury that an accused person has committed a criminal act, but was insane at the time, a special verdict to that effect shall be returned, the court shall then order the accused to be kept in custody as a criminal lunatic till his Majesty's pleasure be known. The prisoner is transferred to a criminal lunatic asylum, and is not set at liberty unless in the opinion of the Secretary of State his liberation is consistent with public safety.

The Medical Superintendent gives the following statistics regarding the 3,220 persons admitted up to the end of the year 1912 into the Broadmoor Asylum for criminal lunatics -

	Total Number Admitted	Admitted for Murder or attempted Murder
Certified to be insane while awaiting trial or judgment,	194	135
Found insane on arraignment,	527	347
Acquitted on the ground of insanity, or found guilty but insane in the terms of the Trial of Lunatics Act, 1883,	1,228	1,061
Reprieved on the ground of insanity	54	54
Found to be insane whilst undergoing sentence,	1,217	168
	<hr/> 3,220	<hr/> 1,765

It will thus be seen that of the 1,765 persons admitted for the crime of murder or attempted murder, less than 28 per cent were found insane on arraignment or before trial, whilst 72 per cent were considered sufficiently sane to be tried.

The defective procedure is productive of evils in a variety of ways. The accused is rarely examined by an expert until a considerable time after the commission of the act for which he was arrested, the law demands proof of

insanity at the time the act was committed, therefore the nearer to it an examination into the state of the prisoner's mind is made, the more likely is it to remove doubts. Again, the less obvious forms of insanity are not unfrequently urged for the first time after the prisoner is condemned, and evidence as to his sanity or insanity may be given by any medical man who happens to have formed an opinion on the subject, this fosters a dangerous tendency to push expertism too far, and, with the aid of the prisoner's counsel, to strive to carry the point at all costs. Much stress is usually laid on real or assumed heredity, the importance of which may be over-estimated, the question is forcibly put by Bucknill<sup>1</sup> when discussing the pleas of insanity in the case of Guiteau, the assassin of General Garfield, President of the United States. "The argument in favour of insanity founded upon the supposed transmission of an hereditary tendency to mental disease has of late been used to most absurd and unjustifiable excess, and I do not know that the interests of justice would be damaged if it were to be excluded altogether in judicial enquiries, for if it could be clearly shown that both a man's parents, and all four of his grandparents, and all his uncles and aunts had been unquestionably insane it would afford no proof whatever that the man himself had been insane. Such evidence would at the most strengthen the presumption that he had been so under circumstances which would otherwise be more doubtful."

Moral imbecility is invariably a stumbling-block to lawyers, who usually regard it as the medical definition of unmitigated depravity. From what has already been said on the subject, it is to be admitted that there is some excuse for this view, inasmuch as the two conditions are not capable of precise distinction. There exists what Maudsley terms "a borderland between crime and insanity, near one boundary of which we meet with something of madness, but more of sin, and near the other boundary of which something of sin but more of madness."

It was asserted in an early section on this subject that delusions are indicative of insanity, it is none the less true that insanity may exist without delusions. This is a hard thing for lawyers to accept, and many of them decline to accept it. To establish the plea in cases of simple moral imbecility, the past family and personal history of the accused must be relied on to a great extent, and in those cases in which the absence of moral perception is due to the onset of a progressive mental disease— as in general paralysis— the concomitant symptoms must be sought for.

The case of *Reg v Edmunds* (C C C, 1872) is a good example of moral imbecility leading to perpetration of criminal acts, by which the lives of innocent people are sacrificed without scruple or remorse in order to avert suspicion from the criminal, and to direct it against those who have done no wrong. Christina Edmunds, aged forty three, was charged with the wilful murder of a little boy named Barker. The boy ate some chocolate creams, which were bought at a respectable confectioner's shop, half an hour afterwards he died with symptoms of poisoning by strychnine, the presence of the alkaloid being subsequently detected in the contents of the stomach. It was proved that the prisoner had obtained a considerable amount of strychnine under false pretences, had got possession of the druggist's poison book, and had torn out leaves which recorded the purchase. It appeared that she had incorporated part of the poison with some chocolate creams, and then asked a small boy to purchase some more creams for her, when he brought them she said they were too large, and sent them back to be changed. Unknown to the boy she substituted the poisoned creams, which, when returned to the confectioner, were placed with his ordinary stock to be sold in due course. One or more of these poisoned sweets caused the death of the boy Barker, who was totally unknown to the prisoner; she also distributed poisoned sweets to many children, who became ill. At the inquest which was held on the body of the deceased

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<sup>1</sup> *Brain*, 1883

she volunteered evidence before being suspected of the crime in order to implicate the confectioner who had sold the sweets, she also wrote anonymous letters to the father of the deceased inciting him to take legal proceedings against the confectioner. This was not done through malice towards the man, but to divert suspicion from herself. She had previously been accused of endeavouring to poison a lady, for whose husband she had conceived a regard, and the whole of this elaborately carried out public poisoning was apparently the result of a scheme to make it evident that the lady's indisposition was also due to poisoned sweets owing to carelessness of the confectioner. It was proved in evidence that the prisoner's father on two occasions had been under restraint, and that he died in an asylum, that one brother had epilepsy and died in Earlewood Asylum, that a sister was hysterical, and had tried to throw herself out of a window, and that other members of the family had suffered from various psychoses. Expert physicians were called to prove that the prisoner was morally insane—she was without intellectual defect and was free from delusions, but she was indifferent to her position and to the enormity of her crimes. She was found guilty and was sentenced to death, but the sentence was subsequently changed to detention in Broadmoor Asylum.

This is a typical case of moral imbecility, all the more so because of its marked resemblance to unbridled depravity. One of the medical witnesses spoke of her being on the borderland between disease and vice, there can be little doubt that mental disease, in the form of imperfectly evolved higher centres, deprived her of the self-control of a sane person, and was the true cause of her criminal conduct.

**Paranoia** appeals more cogently to the legal mind than moral imbecility, but even when delusions are present, it is demanded that they shall be of such a nature as to take away the power of distinguishing between right and wrong. Lawyers attach much importance to the presence of delusions as a sign of insanity, and admit that they may be so dominant as to disturb the judgment to a degree inconsistent with sane conduct. Martin, B., when summing up in the case of *Reg v Townley* (Derby Winter Assize, 1863), said that "what the law meant by an insane man, was a man who acted under a delusion and supposed a state of things to exist which did not exist, and acted thereupon." Thus mere delusions, whatever proof of mental disease they may afford to medical men, afford no proof in the eyes of the law unless the individual is thereby rendered unconscious of the nature and quality of the act of which he has been guilty. Delusions of a lesser type are called partial delusions by lawyers and are not regarded as doing away with the responsibility of a criminal act.

Delusions which frequently bring the subjects of them within the grasp of the law are those of persecution, or of a sense of wrong or injury inflicted that imperatively demands justice. It is to a case of this kind that we owe the legal test of insanity previously mentioned. In the year 1843 a man named M'Naughton shot Mr Drummond, whom he believed to be conspiring with others against his life and character. M'Naughton was acquitted on the ground of insanity, and as there was a general outcry at the supposed failure of justice, the House of Lords propounded certain questions to the judges in order to elicit an authoritative ruling with regard to the plea of insanity. The answers to these questions constitute the law on the subject, as explained in the preceding pages.

Illustrative of the loss of self-control caused by a delusion is the case of *Reg v Dodwell* (CCC, 1878).

The prisoner was a clergyman who became involved in legal proceedings, and after quarrelling with his legal adviser, conducted his own case in such an irregular manner that he did not obtain what he desired. On the strength of this he conceived that he had a grievance against the Master of the Rolls. One morning he awaited the arrival of his Lordship and fired a pistol at him, no injury was inflicted, as the pistol was only loaded with

powder and wadding, the prisoner declaring that his sole object was to direct public attention to his wrongs. At the trial the Master of the Rolls stated that the prisoner was incoherent and irritable, and that he appeared to be under a delusion, no medical evidence was called on either side, and the jury returned a verdict of "not guilty on the ground of insanity." In this case there was no legal evidence of insanity, the prisoner's action might have been due to irascibility and culpable neglect to curb a violent temper, all the evidence went to prove that he knew perfectly well the quality of the act and that he was doing wrong. Possibly the heinousness of the offence—firing at a high legal functionary—was regarded as sufficient to constitute proof of insanity. Dodwell's subsequent history is not without interest in relation to the recurrence of insane impulses, for, although he did not come up to the legal test of insanity, he was none the less a victim of delusional insanity judged from the medical standpoint. After the trial he was removed to Broadmoor, and in 1882 he committed a murderous assault on the chief physician of the asylum by dealing him a heavy blow on the crown of the head with a stone slung in a handkerchief. The motive which instigated him was identical with that which prompted him to fire the pistol at the Master of the Rolls—he stated that as the previous act had not proved sufficient to redress his wrongs, he made up his mind to commit some still more serious act, and had come to the conclusion that nothing less than murder would be sufficient to deliver him from the conspiracy of which he imagined himself the victim.

In relation to the duties of medical men when called upon to pronounce as to the sanity or otherwise of a prisoner, it is to be observed that in many cases it is impossible to arrive at a reliable conclusion without having the individual under observation for some time. Usually several interviews are allowed, for which, and for obtaining the family and personal history of the case, every facility is afforded—but in some cases more is required, which is not obtainable. It should be possible to have such cases placed under skilled care, not only of expert physicians, but of attendants also, who are accustomed to the insane, by close and continuous observation a trustworthy opinion might be formed as to the real nature of the case. No medical man is justified in going into the witness box and (if allowed) delivering himself of dogmatic statements as to the insanity of a prisoner after only a casual interview, unless indeed the mental state is so obvious as not to admit of doubt.

In every respect it would be advantageous if the plea of insanity was disposed of before the trial. The present state of the law is a stumbling-block to this proposal—as interpreted by the judges it will not allow what the Secretary of State is subsequently obliged to concede to public opinion. This leads to those unseemly discussions in the daily papers which so frequently follow a death-sentence. In one class of cases the outcry is the result of a genuine conviction that a judicial blunder has been made, in such cases the end doubtless justifies the means, the result is, that if there is reasonable ground for doubting the convict's sanity, he is sent to Broadmoor instead of being hanged.

An able address on Insanity in relation to offences against the Criminal law, delivered by Orange at the annual meeting of the Medico-psychological Association, with the discussion thereon, is contained in the *Journal of Mental Science* for 1884. The same authority deals exhaustively with the question of Criminal Responsibility in Tuke's *Dictionary of Psychological Medicine*, 1892. See also "Criminal Responsibility," by Charles Mercier, M.B., 1905. The editor has discussed the question in *The Practitioner*, December, 1920.

### **Drunkenness in Relation to Criminal Responsibility.**

The law lays down no definite rules with regard to the plea of drunkenness as a bar to, or in mitigation of, punishment for crime committed under its influence. The principles which usually guide the judges when dealing with crimes of magnitude are—that simple intoxication affords no excuse for the

commission of crime, but if by prolonged drinking the mind is impaired, the condition is regarded as being on a par with ordinary insanity, and may be pleaded as a defence against criminal responsibility. The view taken is—that simple intoxication cannot be held to excuse an offence committed whilst under its influence, because the loss of control was produced by the drunken man's own default. In the case of *Reg v Williams* (Old Bailey, 1886), Denman, J., ruled as follows—that a crime committed during drunkenness was as much a crime as if it were committed during sobriety, and that the jury had nothing to do with the fact that the man was drunk. The prisoner was supposed to know the effect of drink, and if he took away his senses by means of drink, it was no excuse at all. Sir Henry James<sup>1</sup> expresses the opinion that the man who chooses to drink to excess, and, when drunk, from time to time commits acts of brutal violence, must be taught that he is answerable both for being under the influence of alcohol and for the acts such influence induces. The question of intention, however, should it form an essential quality of the offence committed, may be taken into consideration. In the case of *R v Doherty* (16 Cox, C C 306), in which the accused, whilst under the influence of drink, had shot a man, Fitzjames Stephen said to the jury—“Although you cannot take drunkenness as any excuse for crime, yet when the crime is such that the intention of the party committing it is one of its constituent elements, you may look at the fact that a man was in drink in considering whether he formed the intention necessary to constitute the crime. A drunken man may form an intention to kill another, but if he did form that intention, although a drunken intention, he is just as much guilty of murder as if he had been sober.”

The fact that drink does not always affect people to a like degree constitutes a great difficulty. A man either from natural or acquired susceptibility may become mamacal under the influence of an amount of drink that would but slightly affect an ordinary man, such a man under the influence of drink is much nearer the condition of true insanity than that of outrageous drunkenness. The reply obviously is—that a man so constituted should not take drink, and that if he does, it is at his peril. Morally, the question is hard of solution, but the administrators of the law act in accordance with the view just enunciated, and punish the drunkard for committing a crime when he is partly drunk and partly insane, as though he was wholly the former. They are justified in doing so by the evil which would result if those who drink to excess were encouraged to believe that crimes committed under the influence of alcohol would be lightly dealt with.

When mental aberration is due to the remote effects of alcohol, as in **delirium tremens**, judges usually allow that the condition is one of insanity, and it is dealt with accordingly, but even then the legal test for insanity may be applied, and the prisoner's criminal responsibility estimated by it. Great stress is laid by many judges on the permanency of the mental disorder which accompanies delirium tremens, but the exact meaning they attach to the term “permanent” is doubtful. Presumably, it refers rather to a continuance of the symptoms for a definite time after the individual has ceased to indulge in alcohol, than to an absolutely permanent insane condition.

In *Reg v M'Gowan* (Manchester Assizes, 1878), Manisty, J., ruled that “a state of disease brought about by a person's own act—as delirium tremens, caused by excessive drinking—was no excuse for committing a crime unless the disease so produced was permanent.” Subsequently a much less restrictive ruling was given in the case of *Reg v*

<sup>1</sup> *The Times* Newspaper, Jan 4, 1892

*Baines* (Lancaster Assizes, 1886) by Day J., who said "that the question was whether there was insanity or not, that it was immaterial whether it was caused by the person himself or by the vices of his ancestors and that it was immaterial whether the insanity was permanent or temporary. Mr Justice Day further ruled—"that if a man were in such a state of intoxication that he did not know the nature of his act, or that his act was wrongful, his act would be excusable." This ruling is in marked contrast with that previously quoted as given by Denman, J., and also with that given by Bramwell, B., in *Reg v Burns* (Liverpool Assizes, 1865), "that drunkenness was no excuse, and that a prisoner cannot by drinking qualify himself for the perpetration of crime, but if through drunk his mind had become substantially impaired, a ground of acquittal would then fairly arise." The term "substantially" in this ruling replaces the term "permanent" in the ruling given in *Reg v M'Gowan*, above quoted, and seems to point to an impaired mental condition, which, although caused by drinking alcohol to excess, is not due to the immediate presence of alcohol in the system, but to more lasting changes produced in the higher levels of the brain, by which the power of self control is lowered or entirely lost, and the individual is thereby rendered subject to delusions, and is deprived of the knowledge of the nature and quality of his actions.

The most recent case in which drunkenness was pleaded in a criminal charge was that of *R v Beard*<sup>1</sup> in 1919-1920. Beard, who had been drinking, killed a child of thirteen by suffocating her in his attempts to prevent her screaming while he was raping her. At the trial, it was claimed for the defence that he was so drunk that he did not know what he was doing, but he was, nevertheless, found guilty of murder. On appeal the Court of Criminal Appeal quashed the conviction for murder, and found instead a verdict of manslaughter. The case was then referred to the House of Lords, which reversed the decision of the Court of Appeal and upheld the original verdict of murder. In this case there was no doubt that the man was not too drunk to form the intention of committing the crime of rape, and he could not, therefore, be regarded as not responsible for a further act of violence which caused death and was the consequence of the initial crime.

### (b) LUNACY CERTIFICATES.

Before a person of unsound mind can be legally placed under restraint, certain conditions specified by Act of Parliament must be fulfilled. The law jealously safeguards the liberty of the subject, and imposes a number of stringent regulations upon those to whom power is given of placing and receiving insane persons in institutions, or private houses, for the purpose of treatment and of preventing them doing injury to themselves and to others. The Acts which govern these matters are the Lunacy Act, 1890 (53 Vict. ch. 5), and the Lunacy Act, 1891 (54 and 55 Vict. ch. 65). The subject, being of the highest importance to medical practitioners, demands careful and detailed consideration.

Lunatics may be placed under restraint by the following modes of procedure which are varied to suit the exigencies of each individual case:

- Reception Order on Petition
- Urgency Order
- Order after Inquisition
- Summary Reception Order
- Order for Lunatics Wandering at Large, and for Pauper Lunatics
- Reception Order by two Commissioners

**Reception Order on Petition.**—This is the usual mode of procedure in the case of private patients.

The **order** for the reception of the patient is to be obtained by private

<sup>1</sup> *Criminal Appeal Cases*, Part 5, p. 110

application from a "Judicial Authority"—that is to say, either a specially appointed justice of the peace, judge of county courts, or magistrate, lists of judicial authorities are published annually, and may be obtained from the Clerk to the Court of Quarter Sessions for the County, or the Clerk to the Justices. A **petition** for the order must be presented to the Judicial Authority, if possible by the husband or wife, or by a relative of the alleged lunatic, if by another person the reason for this departure must be explained. No person may present a petition unless he is at least twenty-one years of age, and has seen the alleged lunatic **within fourteen days** before its presentation. The petition must be accompanied by a **statement of particulars**, and by **two medical certificates**. Printed forms for all these documents are to be obtained. The petition is considered in private, no one except the petitioner, the alleged lunatic and any one person appointed by him, together with the persons signing the medical certificates, may, without the leave of the Judicial Authority, be present. With the exception of the alleged lunatic and the person appointed by him, all these persons, including the Judicial Authority, are bound to keep the matter secret except when required to divulge the same by lawful authority.

The Judicial Authority if satisfied may make the order forthwith without seeing the patient, or he may appoint a time not more than seven days after the presentation of the petition for enquiries and consideration, he may also visit the alleged lunatic. At the time appointed for the consideration of the petition, he may either make an order or may adjourn the same for any period not exceeding fourteen days, and if he thinks fit he may summon further witnesses. If the Judicial Authority dismisses the petition, he must give the petitioner a statement in writing containing his reasons for doing so. A reception order ceases to be valid after the expiration of seven clear days from its date, except when suspended by a medical certificate of unfitness of lunatic for removal, in which case the lunatic may be received within three days after the date of a medical certificate to the effect that he is fit to be removed.

**Urgency Orders.** In cases of urgency, where it is expedient that the alleged lunatic (not a pauper) shall forthwith be placed under care and treatment, he may be temporarily received *without the intervention of a Judicial Authority*, upon an **urgency order**, accompanied by a **statement of particulars**, and **one medical certificate**. If possible, the urgency order should be made by the husband or wife, or by a relative of the alleged lunatic, it may be signed either before or after the medical certificate. It may be made as well after as before a petition for a reception order has been presented, if made before, it must be referred to in the petition, if made after, a copy of it must be forthwith sent by the petitioner to the Judicial Authority. If the urgency order is not signed by one of the persons named, it must contain a statement of the reasons why it is not so signed, and of the connection of the person signing it with the alleged lunatic. No person may sign an urgency order unless he is at least twenty-one years of age and has seen the alleged lunatic **within two days** before the date of the order. An urgency order remains in force for seven days from its date, or, if a petition for a reception order is pending, until the petition is finally disposed of. The medical certificate must contain a statement that it is expedient for the alleged lunatic to be forthwith placed under care, with the reasons for such statement. The certifying medical practitioner must have personally examined the alleged lunatic not more than **two clear days** before his reception.

**Orders after Inquisition.**—A lunatic found so by inquisition (the procedure will be explained subsequently) may be received upon an order signed by the

committee of the person of the lunatic, or upon an order signed by a Master in Lunacy

**Summary Reception Orders.**—Every constable, relieving officer, and overseer of a parish, who has knowledge that any person within his district or parish, who is not a pauper and not wandering at large, is deemed to be a lunatic, and is not under proper care and control, or is cruelly treated or neglected by those in charge of him, shall within three days give information on oath to a justice being a Judicial Authority under the Act. Such justice may visit the alleged lunatic, but whether he does so or not, he shall direct two medical practitioners to examine and certify as to his mental state, and shall then proceed as if a petition for a reception order had been presented to him. A lunatic as to whom a summary reception order has been made may be taken care of by a relation, or friend, with the consent of the Justice who makes the order, or of the visitors of the asylum in which the lunatic is, or is intended to be, placed

**Orders for Lunatics Wandering at Large.**—Every constable and relieving officer and every overseer of a parish who has knowledge that any person (whether a pauper or not) is wandering at large within his district or parish, and deemed to be a lunatic, shall immediately apprehend and take the alleged lunatic before a justice, or the justice, if information on oath is tendered to him, may require such constable, relieving officer, or overseer to apprehend and bring the alleged lunatic before him. The justice shall then call in a medical practitioner, and if he signs a certificate and the justice is satisfied that the alleged lunatic is a lunatic, and is a proper person to be detained, he may issue an order to that effect. If the medical practitioner certifies in writing that the lunatic is not in a fit state to be removed, the removal order shall be suspended until his fitness is certified to. If a constable, relieving officer, or overseer is satisfied that it is necessary for the public safety, or the welfare of the alleged lunatic, that he should be placed under control before the above-mentioned proceedings can be taken, he may be removed to the workhouse of the union in which he is, and detained not longer than three days, before the expiration of that time the proceedings required by the Lunacy Act shall be taken

**Reception Orders by Two Commissioners.**—Any two or more Commissioners in Lunacy may visit a pauper lunatic or alleged lunatic not in an institution for lunatics or a workhouse, they may call in a medical practitioner, and if he certifies with regard to the lunatic, and they are satisfied that the pauper is a lunatic, they may order his removal to an institution for lunatics

With the exception of the two last named, these regulations do not apply to pauper lunatics or alleged lunatics. The usual proceeding in the case of **pauper lunatics** is for the medical officer of the union to give notice in writing to the relieving officer, or if there be none, to the overseer of the parish where the pauper resides, that a pauper resident within the district is a lunatic and a proper person to be sent to an asylum, this he is bound to do within three days after obtaining knowledge of such pauper lunatic. Every relieving officer, or if there be none, overseer of the parish in which the lunatic resides, who has knowledge either by notice of the medical officer, or otherwise, that a pauper resident within the district or parish is deemed to be a lunatic, shall within three days give notice to a justice having jurisdiction in the place where the pauper resides. The justice then orders the pauper to be brought before him and calls in a medical practitioner to examine him, and if the medical practitioner signs a certificate and the justice is satisfied that the alleged lunatic is a lunatic and a proper person to be detained, he orders his removal to an institution



for lunatics. Thus it will be seen that in the case of **pauper lunatics** only **one** medical certificate is required.

If any lunatic who is detained escapes, he may be retaken at any time within fourteen days without a fresh order. A reception order remains in force for successive periods of one, two, and three years, and afterwards for successive periods of five years. At the end of each period respectively, it may be continued by the Commissioners in Lunacy, on certification by the medical officer of the institution, or by the medical attendant of a single patient, that the patient is still of unsound mind and is a proper person to be detained under care and treatment. A patient may be discharged on the direction of the petitioner for the reception order, or, if the petitioner is dead or incapable, by the nearest of kin, or by the Commissioners. But if the medical man in charge certifies that the patient is dangerous and unfit to be at large, he shall not be discharged unless two of the visitors of the asylum, or the Commissioners visiting the hospital or house, consent in writing to the patient's discharge. Two Commissioners, one of whom shall be a medical and the other a legal Commissioner, may order the discharge of any patient.

The principle which governs the provisions of this Act is that no person can be legally placed under restraint as a lunatic without an order obtained from a Judicial Authority. Exceptions to this principle are constituted by cases which are dealt with by inquisition, and by two Commissioners. Urgency orders are made without the intervention of a Judicial Authority, but they are only temporary and provisional, being intended to prevent harm happening to or by the lunatic until the necessary formalities have been observed to permanently place him under restraint, so that although an alleged lunatic has been admitted into an institution for lunatics on an Urgency Order the entire procedure relating to obtaining a reception order on petition has to be gone through just as though the alleged lunatic had not been previously dealt with.

### Medical Certificates in Lunacy.

Apart from lunatics found so by inquisition, lunatics wandering at large, pauper lunatics, and lunatics who are temporarily taken care of on an Urgency Order, **two** medical certificates are required before any person can be placed under restraint as a lunatic.

Every medical certificate must state the facts on which the certifier has formed his opinion that the person to whom the certificate refers is insane, he must distinguish facts observed by himself from facts communicated by others, and it is important to note that a reception order will not be made upon a certificate founded only on facts communicated by others. In respect to **orders on petition** each medical practitioner who signs a certificate must have personally examined the alleged lunatic **within seven clear days** before the date of the presentation of the petition, and in all other cases within seven clear days before the date of the order. Where two medical certificates are required, each medical practitioner must examine the alleged lunatic separately from the other. In the case of **urgency orders** the practitioner must have examined the alleged lunatic **not more than two clear days** before his reception. A medical certificate may not be signed by the petitioner for an order, nor by the person signing the urgency order, nor by any near relative, partner, or assistant of the petitioner or person. One of the medical certificates accompanying a petition for a reception order should be under the hand of the usual medical attendant of the alleged lunatic, if this is not practicable the reason

must be stated in writing by the petitioner to the Judicial Authority to whom the petition is presented. No certificates shall be signed by persons interested in the institution or house to which the lunatic is going, nor by any near relations of such persons, nor by two medical men related to or in partnership with each other, or standing in the relation of principal and assistant.

A medical practitioner who has signed a certificate upon which a reception order has been made shall not be the regular professional attendant of the patient while detained under the order. A person for whom a reception order on petition has been obtained may be placed in a "single-patient house"—that is, a house in which one lunatic only is detained on payment exactly as in an asylum, or in a licensed house *i.e.*, private asylum. Under special circumstances the Commissioners may allow more than one patient to reside in the same house under the same conditions as if each of them were a single patient. If the usual medical attendant of the lunatic desires to continue in attendance during the detention of the lunatic in a single-patient house, he must not only *not sign either of the certificates*, but neither he, his partner, nor any of his near relatives must derive any profit from the charge of the patient. The Commissioners may direct how often a single patient is to be visited by a medical practitioner, until such direction is given the patient must be visited at least once every two weeks. Any two Commissioners may direct that the medical attendant of a single patient shall cease to act in that capacity, and that some other person shall be employed in his place. The Commissioners may at any time require from the medical attendant of a single patient a report in writing as to the patient, with such particulars as the Commissioners may direct, this is in addition to any periodical reports required by law to be sent to the Commissioners.

The Commissioners may require a report of the mental and bodily condition of a lunatic, or alleged lunatic, who, without order and certificates, is detained or treated as a lunatic by any person receiving no pay for the charge, or who is in any charitable, religious, or other establishment, not being an institution for lunatics, the report is to be furnished by a medical practitioner, and repeated periodically if required. The Commissioners may also visit such patient, and may exercise all the powers (except that of discharge) given to them as to patients in an asylum or as to single patients. If they think fit they may inform the Lord Chancellor, who may make an order for the discharge of such patient, or for his removal to an institute for lunatics. This is a departure from the former custom, which was that the Commissioners had no control over patients treated in private houses where no one derived any pecuniary benefit from their detention.

**Examination of Alleged Lunatics in Relation to Lunacy Certificates.**—The objects of the examination are to determine whether the individual is or is not insane, and if insane whether he is a fit and proper person to be placed under restraint. The distinction is important, as a person may be undoubtedly insane, and yet neither his language nor his actions may justify a medical practitioner in certifying that he ought to be sent to an asylum. The printed forms of certificates have marginal- or foot-notes explanatory of the mode in which they are required to be filled up, a reference to the appended form will at once show what is meant. Any practitioner who is not accustomed to fill up these forms should carefully read over the directions, and be sure he fully understands them before commencing to write. It is to be remembered that the law requires absolutely literal accuracy, the least omission is sufficient to invalidate the whole document.

When examining an alleged lunatic—which is best done in the character of a medical man, although in exceptional cases it may be advisable to personate a casual caller or some one other than a doctor—if his words or actions do not at once reveal insanity the examiner will soonest obtain an opportunity of judging as to his mental state by directing the conversation to personal matters. Should the case be one of delusional insanity some information previously obtained as to the nature of the delusion, or delusions, will materially help the investigation. Delusions may be of two kinds: they may carry their own refutation, or they may present no abstract inconsistency. If, in a certificate, it is stated that the patient says he is the Holy Ghost, comment is superfluous, but if it is stated that he says he is starving for want of money, no delusion is evident unless information is added such as the further statement that he is really a wealthy individual with an income of a thousand a year. The officials into whose hands the certificates come know nothing beyond what is contained in them, about the medical features of the case: the matter, therefore, should be made perfectly clear and understandable. When the alleged lunatic is not subject to delusions there may be considerable difficulty in obtaining anything definite to enter in the certificate, as facts observed by the certifier, in such cases more than one visit, possibly several, may be necessary before sufficient evidence of insanity is forthcoming as to warrant certification.

In relation to facts it may be well to direct attention to what might be supposed to be a very obvious difference, absence of appreciation of which, however, has often led to rejection of certificates, it is the distinction between facts indicative of insanity, observed by the certifier at the time of his visit, and communicated facts. For example, the fact was communicated to a certifier that the alleged lunatic was intemperate, on the strength of this, and seeing him drink a glass of beer, the statement that the man's habits were intemperate was incorrectly incorporated in a certificate under the head of fact observed by the certifier. It is by no means easy in all cases to obtain facts that afford conclusive proof of insanity: before entering facts of doubtful significance, their value should be well weighed by the practitioner. He should ask himself whether they afford reasonable proof of mental disease, and how far he could depend upon them to support his opinion if challenged in a court of law. In many cases of undoubted insanity it is absolutely impossible to obtain any one fact conclusive of the condition, and consequently dependence has to be placed on accumulative evidence. In such instances special care must be taken to reject matter that will not bear the test of legitimate cross-questioning. Clouston gives some ludicrous examples of "facts"—"He is incoherent in his appearance", "Eyes restless and wandering, but following the usual occupation of breaking stones", "Reads his Bible, and is anxious about the salvation of his soul". It is to be remembered that even though the Commissioners accept a certificate, legal proceedings may subsequently be taken by the alleged insane person, and the certifier may be placed in the witness-box and cross-examined on the statement he has made.

Mere defect of intelligence, which may exist without delusions, is another condition in which it is difficult to obtain evidence of sufficient value as to be suitable for a certificate of lunacy. Tests of mental capacity in such cases should be restricted to those which are compatible with the station in life of a person under examination, a want of intelligence that would be abnormal in a man possessing all the advantages of a good education would be much less significant in the case of an agricultural labourer. A degree of mental

## LUNACY ACT, 1890

## CERTIFICATE OF MEDICAL PRACTITIONER

IN THE MATTER OF *A B* of <sup>1</sup> in the County <sup>2</sup>  
<sup>3</sup> an alleged lunatic

I, THE UNDERSIGNED, *C D*, do hereby certify as follows —

1 I am a person registered under the Medical Act, 1858, and I am in the actual practice of the medical profession

2 On the day of 19, at <sup>4</sup> in the county <sup>5</sup>  
 of (separately from any other practitioner) <sup>6</sup> I personally examined the said *A B*, and came to the conclusion that he is a (lunatic, an idiot, or a person of unsound mind) and a proper person to be taken charge of and detained under care and treatment

3 I formed this conclusion on the following grounds, viz —

(a) Facts indicating insanity observed by myself at the time of examination, viz —<sup>7</sup>

(b) Facts communicated by others, viz —<sup>8</sup>

*(If an urgency certificate is required it must be added here)*

4 The said *A B* appeared to me to be (or not to be) in a fit condition of bodily health to be removed to an asylum, hospital, or licensed house <sup>9</sup>

5 I give this certificate having first read the section of the Act of Parliament printed below

Signed

<sup>10</sup> of

Dated,

*Extract from Section 317 of the Lunacy Act, 1890*

Any person who makes a wilful mis statement of any material facts in any medical or other certificate, or in any statement or report of bodily or mental condition under this Act, shall be guilty of a misdemeanour

<sup>1</sup> Insert the residence of patient

<sup>2</sup> City or borough, as the case may be

<sup>3</sup> Insert profession or occupation, if any

<sup>4</sup> Insert the place of examination, giving the name of the street, with number or name of the house, or should there be no number, the Christian and surname of occupier

<sup>5</sup> City or borough, as the case may be

<sup>6</sup> Omit this where only one certificate is required

<sup>7</sup> If the same or other facts were observed previous to the time of the examination, the certifier is at liberty to subjoin them in a separate paragraph

<sup>8</sup> The names and Christian names (if known) of informants to be given, with their addresses and descriptions

<sup>9</sup> Strike out this clause in case of a private patient whose removal is not proposed

<sup>10</sup> Insert full postal address

weakness that may be sufficient to justify a man being deprived of the management of his monetary affairs, may be insufficient to justify his being deprived of liberty of action in other respects, to warrant this something more is needed than absence of a certain degree of intelligence. On more than one occasion, when giving evidence on the subject of criminal responsibility, psychological experts have stated in the witness-box that although, on the ground of mental disease, they held the prisoner to be irresponsible for the crime he had committed, they would not have seen their way to certify him as a fit and proper person to be placed under restraint.

Actions and utterances must also be considered in relation to the social position and previous habits of the individual. It is deplorable for a costermonger to use bad language to his wife or to assault her and destroy the furniture in his house, but such proceedings are of much less weight as indications of insanity than when the person implicated is, or rather has been, a sedate clergyman. General expressions, as "excited and wild-looking," should, as far as possible, be avoided, and something more precise substituted. In recording utterances the actual words used by the alleged lunatic should be given.

A medical man is supposed by the law to exercise his own judgment when deciding as to the sanity or insanity of a patient, he must be on his guard not to be unduly influenced for or against the statements of the wife, husband, or other members of the household. In some instances it is to the interest of the relatives to further the impression of the patient's sanity, and they will consequently seek to minimise the significance of his actions, attributing any out-of-the-way conduct to mere eccentricity, or to family worries, if the object is the converse, every action is unfavourably interpreted, and the truth is by no means strictly adhered to.

**The Legal Responsibility of Medical Practitioners in Relation to Certificates of Lunacy.**—No medical practitioner is bound to certify, but if he undertakes to do so he is responsible for any breach of duty. A medical man who makes a wilful mis-statement of any material fact in a medical certificate is guilty of a misdemeanour. Up to the time of the Lunacy Act of 1889, medical men who signed certificates of lunacy were liable to be the victims of harassing legal proceedings, although they acted in perfect good faith. Nothing can exceed the vindictiveness of some persons who, having been insane, have sufficiently recovered as to be released from restraint, they acquire a fixed idea that they have been the victims of a vile conspiracy, and display a most objectionable pertinacity in attacking (through the law courts) each and every person concerned in the case. The medical practitioners who certified are invariably selected, and as a rule they fare badly, especially if they have committed any technical error. The last Act has a clause (section 330, re-enacting section 12 of the Act of 1889), which provides for vexatious proceedings directed against those who have been engaged in carrying out the Act, and medical practitioners come in for a share of the protection thus conferred. The gist of the section is that such proceedings may, "upon summary application to the High Court, or a judge thereof, be stayed upon such terms as to costs and otherwise as the Court or judge may think fit, if the Court or judge is satisfied that there is no reasonable ground for alleging want of good faith and reasonable care."

This section renders it incumbent on the medical practitioner who is proceeded against to satisfy the Court that there is no reasonable ground for alleging want of good faith or reasonable care. "Good faith" is to be interpreted in its common-sense meaning—that is, if the transaction was honestly carried out and there was no attempt to abuse the power conferred by law.

it was done in good faith. "Reasonable care" is held to have the same significance that it has in ordinary cases of malapraxis: the employment of such an amount of care and skill as a medical practitioner may reasonably be expected to possess.

The first case tried under this section was that of *Toogood v Wilkes* (Queen's Bench Division, 1889). The plaintiff brought the action against a medical practitioner, to recover damages "for injury to the plaintiff from the defendant's negligence as a medical man, and for damages for injury to the plaintiff by reason of the defendant having negligently and wrongfully signed a certificate of the plaintiff's insanity whereby he was detained in a lunatic asylum." The judge, Field, J., ruled that the onus lay with the defendant to satisfy him that there was no reasonable ground for alleging want of good faith or reasonable care, and after hearing the arguments, held that the defendant had made out that there was no want of good faith or reasonable care on his part in signing the certificate, the action was therefore stayed. This is a step in advance when compared with the former state of things, although it seems strange that the defendant was not allowed costs, the judge declining to give them unless the plaintiff appealed, which he omitted to do. Another case of a similar kind, *Mason v Marshall Shaw, and Gauchard* (Bristol Spring Ass., 1888), in which the contention was that the medical men who had signed the certificates had not exercised sufficient care, and that the certificates were criminally weak, was tried before the Act containing the relief section previously quoted was passed. The judge left the following questions to the jury:—(1) Did the doctors sign the certificates negligently and without due care? (2) Were the conduct, behaviour, and appearance of the plaintiff such as to induce the defendants to believe that she was a person of unsound mind, and a proper person to be taken care of and detained under treatment, and were the acts complained of done by the defendants honestly acting upon such belief? (3) Was the plaintiff at the times in question of unsound mind and a proper person to be taken care of and detained under treatment? The jury answered all the questions in favour of the defendants. The section of the Lunacy Act relating to the staying of vexatious proceedings affords very necessary protection to medical men who have to deal with cases of alcoholism followed by insanity, which may be of a fugitive type. In one such instance, *Williams v Beaumont and Drake* (Court of Appeal, 1894), in which the plaintiff appealed against the decision of the Divisional Court staying an action entered by him against the defendants, it was proved that plaintiff was received into the workhouse whilst suffering from alcoholism, followed by melancholia with suicidal tendency. The defendants, who were the medical officers of the workhouse, decided it best to send the plaintiff to the asylum, and accordingly certified him. On admission into the asylum no indications of insanity were observed, and, after having been kept under observation for a few weeks, he was discharged, hence the proceedings. The Master of the Rolls, with Lord Justice Kay and Lord Justice Smith, in dismissing the appeal, said that they saw no reason for suggesting that the defendants had not acted with reasonable care and in good faith. Lord Justice Smith added that it is immaterial what the plaintiff sets out in the statement of claim, such as trespass, for example, if the action is a proceeding taken against a doctor for something done in pursuance of the Act, the whole question resolves itself into one of reasonable ground for alleging want of good faith or reasonable care.

These cases illustrate the importance attached by the judges to the exercise of such an amount of skill and care as a medical practitioner might be reasonably expected to bring to bear in a case of the kind; it is, further, of equal importance that the spirit as well as the letter of the law should be fulfilled, any shortcoming in this respect is liable to be severely dealt with. In the case of *Weldon v Winslow* (Nisi prius, 1884) the defendant's conduct was held to be irregular and not to conform with the mind of the law.

The defendant, a medical man, was asked by the plaintiff's husband to see his wife and give an opinion as to her sanity; he did so, and reported that she was insane, advising removal to an asylum, and it was arranged that the lady should go to the defendant's own private asylum. This was an undoubted mistake and a violation of the spirit if not of the letter of the law, which provides that persons who are interested in the institution to which a patient is going shall not sign either of the certificates. Although the defendant did not actually sign a certificate he sent two other medical men (after having himself determined the question), and in this way violated the spirit of the law. A further irregularity was

committed by the two medical men who went to examine the patient instead of visiting her separately, as required by law, they went together, one waiting outside the room while the other conducted his examination. In commenting on these proceedings the judge, Mr Justice Manisty, said that if the defendant "had it at all in his mind that the plaintiff should be taken to his asylum, he ought to have told Mr Weldon at once, 'I can take no part in these proceedings. I can take no part in obtaining certificates or getting the order. You must get some one else to act in the matter.'" The judge also asked whether the way in which the examination was carried out satisfied the law that the medical men should be absolutely independent, and should each exercise an independent judgment.

### (c) JUDICIAL INQUISITION AS TO LUNACY.

The Judge in Lunacy may, upon application by order, direct an inquisition whether a person is of unsound mind and is incapable of managing himself and his affairs. The inquisition may take place before a jury if the alleged lunatic demands one, unless the Judge in Lunacy, after personal examination of the alleged lunatic, is satisfied that he is not mentally competent to form and express a wish to that effect (53 Vict ch 5, sec 90).

The object of an **inquisition** (*de lunatico inquirendo*) or **commission of enquiry** is to ascertain whether a certain person is, or is not, fit to retain charge of his affairs, and whether he ought, or ought not, to be placed under restraint. It is to be clearly understood that the two propositions are separate and distinct: the first does not necessarily include the second, although in this case the second includes the first. A man may be found incapable of managing his property by reason of deficiency of intellect, but it does of necessity follow that he is a fit and proper person to be placed under restraint, if the commission of enquiry find that he ought to be detained under care and treatment, then the deprivation of civil rights: the management of his own affairs follows. Lunatics who are taken charge of after inquisition, whether under restraint or not, are called "**Chancery Lunatics**," because the Lord Chancellor is at the head of the administration of the Lunacy Laws, and such lunatics are under his care.

It is to be observed that deprivation of civil rights is not a necessary result of placing a patient under restraint in the ordinary way by order on petition, if competent, such a person can make a will, and if liberated from the asylum, can at once resume the management of his affairs.

To obtain a Commission of Enquiry an application must be made to a Judge in Lunacy by one or more persons interested in the alleged lunatic, supported by affidavits taken by medical practitioners as to his mental condition, if the Judge in Lunacy is satisfied that there is ground for the inquisition, the cause is duly tried. The medical men who have signed certificates, or taken affidavits, with probably others also, representing both sides, are examined and cross-examined on oath as at an ordinary trial. If found to be of unsound mind, and a fit and proper person to be placed under restraint, the lunatic is treated as though found so on petition, and further, his property is vested in the hands of a "committee of estate," which acts as his trustee and manages his affairs. The Lunacy Act limits the enquiry to things said and done by the alleged lunatic within a period of two years previous to the inquisition. As previously mentioned, he may be found mentally incompetent to manage his estate, and yet not incapable of managing himself, in this case his affairs are placed in the hands of a "committee," but he is not deprived of freedom of action in other respects, he can remain in his own house and go about as he pleases.

Commissions of enquiry are usually held in the case of men who, by acts of absurd extravagance, are ruining their estates, the object being to put

an end to this even though the owner of the estate may be not sufficiently insane as to warrant his being detained under restraint. Inquisition is an expensive way of determining the mental condition of a person alleged to be of unsound mind, not so much so, however, as before the passing of the recent Acts.

#### (d) TESTAMENTARY CAPACITY.

By this is meant competency of mind to make a will disposing of personal or real property. In order that a will shall be valid, the law requires a "disposing mind" on the part of the testator. The interpretation placed on this term is somewhat elastic, and often depends upon the nature of the will itself rather than upon the mental condition of the testator at the time he made it. The law is very jealous of any interference with documents of this kind, and even when the testator can be proved to have been of unsound mind, if the will is rational and of such a nature as to be consistent and free from unreasonable prejudice, or from indications of undue influence exercised by others, it is usually upheld. A patient labouring under a delusion is not necessarily thereby incapacitated from making a will. If the delusion is on a subject apart from the provisions of the will, the document itself need not be affected by it, it is quite possible for a man to have delusions of such a nature as not in the least to interfere with his capacity to make a rational will having the same disposing qualities with respect to his relatives as though he was free from any such delusion. A delusion bearing on the provisions of the will is a different matter, and might be held to invalidate the document.

Wills good in law have been made by patients in asylums.

In the case of *Banks v Goodfellow* (Queen's Bench, 1870), an attempt was made to upset the will of the testator, John Banks, who had been in an asylum, and who was subject to delusions. He believed himself to be persecuted by devils, which he stated were visibly present, and he also believed that a man who had been dead for a long time, and to whom he had a great aversion, was still alive, and that this man pursued and molested him. Notwithstanding the existence of these delusions the will, which was made in favour of a niece who had lived with the testator, was upheld by the ruling of four judges sitting in Banco, one of whom, Lord Chief Justice Cockburn, in delivering judgment said—"That the existence of a delusion compatible with the retention of the powers and faculties of the mind will not be sufficient to overthrow the will unless it were such as was calculated to influence the testator in making it."

Wills are not unfrequently disputed on account of alleged undue influence, exercised by some one who has access to a weak-minded person during the latter portion of his life. Unlimited influence can be brought to bear by a designing person on people who are naturally of feeble intellect or who have been rendered so by bodily disease. In this way perfect strangers succeed in obtaining utterly disproportionate legacies, or even the bulk of an estate, to the exclusion of those who have a rightful claim. Few medical men who have seen much practice have not had opportunities of observing the assiduity with which a distant relative, or friend, of an elderly individual who lives alone and is possessed of money but is devoid of sons or daughters, watches and hovers round with ceaseless attention, brings and prepares all sorts of delicacies, and gradually insinuates himself, or more frequently herself, into the very existence of the sick person, until his volition is paralysed, and he yields an unquestioning assent to all suggestions made by the beneficiary *in posse*. In such a case, the will, if made in favour of the attentive person, is very likely to be contested, and it behoves the medical attendant of an invalid thus cajoled to keep his



ears and eyes open, as he is sure to be subpoenaed on one side or other when the case comes before the Probate Court

The condition which may render testamentary capacity doubtful are those due to disease, to feebleness of mind apart from disease, and to the presence of delusions or other indications of unsoundness of mind. A frequent cause of doubt as to testamentary capacity is the occurrence of the stage of lethargy with partial loss of coherence of thought which is the precursor of death. A few questions put to the sick person will probably determine whether the mind is sufficiently clear or not, if he can recapitulate the provisions of the will he is desirous of making, after having once announced them, it is enough to establish his testamentary capacity. The necessary questions may be asked of the dying person in the presence of those who surround his bed—in such a case witnesses are an advantage rather than otherwise. It is different, however, in cases of simple feebleness of intellect occurring in people who are not about to die, the medical practitioner, whose opinion as regards testamentary capacity is sought, should always insist upon speaking to such persons alone. The mere presence in the room of any one who has acquired an influence over their minds, without the utterance of a word, will render the investigation futile. In addition to questions which are intended to test the mental capacity of feeble-minded testators, others should be put with the view of ascertaining whether any undue influence is being exercised over their minds. If delusions are present, their influence on the “disposing mind” of the testator should be estimated as far as possible, he should be asked to state the principal provisions of the will, and if any clause appears contrary to what might be reasonably expected, to explain the motives that influence him in so willing.

The occurrence of **aphasia** may interfere with testamentary capacity, no rule can be formulated for the various forms and degrees of this defect, each case must be judged on its own merits. The primary consideration is—Can any form of aphasia exist without such impairment of the mental powers as to invalidate a will or other legal document executed by a person labouring under this defect? Uncomplicated **motor aphasia** does not necessarily incapacitate a patient from executing such a document, it is obvious that a primal insane condition is excluded, such as dementia with aphasia. In pure motor aphasia the patient can signify by gesture his assent to, or dissent from, any proposition—he is quite competent to make his desires known. Pure motor aphasia is not inconsistent with the accomplishment of mental processes which demand differentiation and combination of ideas. Edmunds<sup>1</sup> records the case of a lady who had right-sided hemiplegia with complete motor aphasia, she could neither speak nor write, but the mind remained intact. The patient made a will by selecting and grouping cards on which were printed the names of persons, of offices such as executorships, and descriptions of property. The solicitor received his instructions from the cards thus selected and made the will, which was proved in the Probate Court. **Sensory aphasia** cannot be disposed of so readily, the very nature of the condition gives rise to doubt as to whether the sufferer comprehends what is said to him, although the mental powers are not necessarily lost. In relation to this question Ross<sup>2</sup> observes that “when a lesion is situated in or near to the sensory inlets, a disorder of language results which is out of all proportion to the general impairment of the reasoning faculties.” Defect of the ordinary channels of communication with the mind of the patient constitutes an essential difficulty, if there is absolutely no channel by which ideas can be conveyed, the individual is obviously incompetent to

<sup>1</sup> *Brit Med Journ.*, 1900.

<sup>2</sup> *Med Chron.*, 1887.

execute a legal document, but if, though imperfect, a channel exists—either the visual or the auditory paths being intact— it would be possible to secure the necessary cognition. Bramwell<sup>1</sup> directs attention to the fact that a moderate degree of word-deafness is more likely to interfere with the will-making capacity of a testator than a much greater degree of word-blindness. Mills,<sup>2</sup> who discusses the medico-legal relations of aphasia at some length, holds that it ought not to be necessary for competency that the person should be responsive by every channel of communication. It is to be borne in mind that an aphasic testator, by reason of the imperfection of his channels of communication, may be unable to grasp the provisions of a lengthy or a complicated will, and yet may be competent to give an intelligent assent to those of a short, straightforward document.

In no case should a medical man allow himself to be influenced by sympathy with the survivors, and countenance the signing of a will, the provisions of which may be just and equitable, by a person who from any cause is incompetent to do so. Attempts are not unfrequently made to complete a will by the almost unconscious assent of a dying person, the document having been drawn up according to his directions at an early stage of the illness when his mind was still active. The practitioner's duty is clearly defined, it is limited to ascertaining the fitness or otherwise of the testator to understand the provisions of the document at the time that he signs it. It is to be remembered that the will may be disputed, and the state of the testator's mind may have to be described on oath under cross-examination.

#### (e) FEIGNED INSANITY.

Insanity is not unfrequently feigned in order to escape punishment for a criminal act. Usually some form is selected that corresponds with the popular idea of insanity, the symptoms imitated being those which demonstrate mental disorder to the most casual observer. Mania, delusional insanity, melancholia, and dementia are generally the types to which the sham lunatic devotes his attention. The cases which come under the notice of the medical jurist are almost invariably those of criminals who are already in the hands of the police. Unless there is a history of previous attacks, the occurrence of symptoms of insanity in a prisoner are naturally suspicious, all the more so if they are of a pronounced character. When **mania** is imitated there is a tendency to take the more excited type, which renders detection much easier than if a lower kind of exaltation was aimed at. The popular idea of mania is limited to what is described as raving madness, and the shammer consequently believes that the more furious his demeanour the more genuine does the attack appear. Fortunately for those who are called upon to decide the question, nature places a limit on the powers of endurance of a sane individual, no man who is not really insane can keep up for long together the constant action and vociferation characteristic of the true acute maniac. Exhaustion of the physical powers, which the real maniac seems to be able to defy, asserts itself, and the impostor falls asleep, and as a rule sleeps soundly, which is in direct contrast with the fitful sleep of acute mania. A true maniac will often pass several consecutive nights and days without sleep, an impostor is utterly unable to do so. If the suspected person manages to do without sleep for a long time, probably a hypodermic injection of morphine, which would produce no effect on a genuine case of mania, will procure sleep, and clear up doubts.

<sup>1</sup> *Brit. Med. Journ.*, 1897.

<sup>2</sup> *Review of Insanity and Nervous Disease*, 1891.

**Melancholia** is not often feigned. In a suspected case the patient must be watched without his knowledge to see if there is any change in demeanour when alone. If he sleeps soundly it is improbable that he is the subject of profound melancholia. The state of the tongue may afford evidence, it is often coated in genuine insanity. If suspicion is strongly aroused, and the individual refuses to speak or take any notice, a strong faradic current applied to various sensitive parts of the skin, by means of a stiff wire-brush, may promote recovery. **Delusions** are usually over-acted by the malingerer, although he may succeed in giving rise to considerable doubt in the mind of the observer as to the real nature of the case. Robertson<sup>1</sup> relates a case of feigned insanity with delusions in a man who was awaiting trial for housebreaking.

The prisoner first drew attention to his assumed mental state by a feigned attempt at suicide by hanging. He then refused to work, would not speak when addressed, and took very little food. Now and then he lay on the floor staring at the ceiling as though he saw some one there. He professed entire ignorance of any circumstances relating to the crime of which he was accused, but declared that he was to be tried for murder, and stated that it was quite true that he had murdered a woman by pushing her into a canal where she was drowned, he also said that the woman came into his cell at night and offered him a razor. He shed tears when telling the story of his avowed crime, and insisted that it was true, although when assured that the woman was alive and well he expressed his thanks, subsequently, however, he recurred to the feigned delusion. Besides this fixed delusion he said that thousands of rats came into his cell at night, and made other absurd statements. At one interview he declared he had committed murder, that there was no hope for him, and that he was eternally lost, at the next, he said that he possessed £400, expected to inherit £4,000, and owned the island of St Helena. This discrepancy was significant, the character of the exaltation was inconsistent with almost simultaneous depression. The attempt was kept up to the commencement of, and during, the trial, but after conviction he confessed that the whole affair was an imposition.

The chief points of distinction are that a lunatic, as a rule, insists that he is sane, whereas a malingerer obviously tries to produce the impression that he is insane. If he is incoherent, his incoherency exceeds that of any lunatic, except, perhaps, that of the delirious maniac. If he has delusions he will show considerable perspicacity in relation to the delusions, but will, at the same time, exhibit the mental state of an imbecile as regards other matters, especially relating to himself and his doings. It has been said that although the sham lunatic may be dirty as far as his surroundings go, he will spare his person. This may be so as a rule, but it is not invariably the case, some will smear their persons with excrement and other filth, having probably seen these actions done by a genuine lunatic.

### CARE OF MENTALLY DEFECTIVE PERSONS.

The Mental Deficiency Act, 1913, provides for the care and treatment of mentally defective persons, and repeals with certain provisos the Idiots Act of 1886. The following are the classes of persons who are deemed to be mentally defective within the meaning of the Act—

(a) **Idiots**; that is, persons so deeply defective in mind from birth or from an early age as to be unable to guard themselves against common physical dangers.

(b) **Imbeciles**; or persons in whose case there exists, from birth or from an early age, mental defectiveness not amounting to idiocy, yet so pronounced that they are incapable of managing themselves or their affairs, or, in the case of children, of being taught to do so.

<sup>1</sup> *Journal of Mental Science*, 1883

(c) **Feeble-minded persons**; that is, persons in whose case there exists, from birth or from an early age, mental defectiveness not amounting to imbecility, yet so pronounced that they require care, supervision, and control for their own protection or for the protection of others, or, in the case of children, that they, by reason of such defectiveness, appear to be permanently incapable of receiving proper benefit from the instruction in ordinary schools

(d) **Moral imbeciles**; or persons who from an early age display some permanent mental defect coupled with strong vicious or criminal propensities on which punishment has had little or no deterrent effect

A person who is a defective may be sent to an institution for defectives or placed under guardianship under the following circumstances —

(a) **At the instance of his parent or guardian**, if he is an idiot or imbecile, or at the instance of his parent if, though not an idiot or imbecile, he is under the age of twenty-one, or

(b) **If, in addition to being a defective, he is a person** who is found neglected, abandoned, or without visible means of support, or cruelly treated, or is guilty of a criminal offence or liable to be sent to an industrial school, or is undergoing imprisonment or detention in a reformatory, industrial school, inebriate reformatory or institution for lunatics, or is an habitual drunkard within the meaning of the Inebriates Acts, or in whose case notice has been given by the local education authority as described later, or who is in receipt of poor relief at the time of giving birth to an illegitimate child or when pregnant of such child

The local education authority is required to give notice of all defective children over the age of seven, who have been ascertained to be incapable, by reason of mental deficiency, of receiving benefit or further benefit in special schools or classes, or who cannot be instructed in a special school or class without detriment to the interests of the other children, or where there are special circumstances rendering it desirable that they should be placed under supervision or guardianship. Notice must also be given of children who, on or before attaining the age of sixteen, are about to be withdrawn from a special school or class, and in whose case the local education authority are of opinion that it would be to their benefit that they should be sent to an institution or placed under guardianship

Where the power to place a defective in an institution or under guardianship is exercised by the **parent or guardian** two medical certificates are required, one of which must be from a medical practitioner approved for the purpose by the local authority or the Board. If the defective is not an idiot or imbecile the certificates must also be signed by a judicial authority for the purposes of the Act, after such enquiry as he shall think fit

A defective subject to be dealt with otherwise than by a parent or guardian may be so dealt with --

(a) Under an order made by a **judicial authority** on a petition presented under the Act, or

(b) Under an order of a **court** in the case of a defective found guilty of a criminal offence

(c) Under an order of the **Secretary of State** in the case of a defective detained in a prison, criminal lunatic asylum, reformatory, etc

**Requirements as to the Making of Orders.**—An order of a judicial authority is obtainable upon a private application by petition made by any relative or friend of the alleged defective, or by any officer of the local authority authorised under the Act in that behalf. The petition must be accompanied with two

medical certificates, one of which must be signed by a medical practitioner approved for the purpose by the local authority or the Board, or a certificate that a medical examination was impracticable, and by a statutory declaration made by the petitioner and by at least one other person (who may be one of the persons who gave a medical certificate), stating the class of defectives to which the person concerned is alleged to belong, and other particulars and attendant circumstances of the case

Upon presentation of the petition and other documents, the judicial authority must either visit the person to whom the petition relates or summon him to appear before him. Proceedings before the judicial authority may, if he thinks fit, and shall, if the person to whom the petition relates so desires, be conducted in private. If the judicial authority is satisfied that the person is a defective, and also that he is subject to be dealt with under the Act, he may, if he thinks it desirable to do so in the interests of the person, make an order sending him to an institution or appointing a suitable person to be his guardian. But where the petition is not presented by the parent or guardian the order shall not be made without the consent in writing of the parent or guardian unless such consent is unreasonably withheld or they cannot be found.

If the judicial authority is not satisfied he may adjourn the case for more than fourteen days for further evidence to be produced, or he may dismiss the petition.

Where an order is made by a court, the Act merely requires the court to be "satisfied on medical evidence" that the person concerned is a defective. Where the Secretary of State acts in the case of a person undergoing imprisonment, etc., two medical certificates are required.

An order remains in force for a year, and then, subject to various safeguards, may be renewed for another year, and thereafter for successive periods of five years.

The central administration of the Act is exercised by the Board of Control, which is now a branch of the Ministry of Health. Local administration is exercised through the County Council or County Borough Council.

### **PLACING HABITUAL DRUNKARDS UNDER RESTRAINT.**

An habitual drunkard may be guilty of actions fully as obnoxious to the well-being of his family and quite as indicative of inability to manage himself or his estate as those committed by a person, who by reason of disease of the mind is a fit and proper person to be placed under restraint, and yet, in such a case (unless the individual is actually insane apart from inebriety), the law does not admit of the procedure applicable to lunatics being put in force. In recognition of this difficulty an Act was passed in 1879 (42 and 43 Vict., ch. 19) to facilitate the control and cure of Habitual Drunkards, the duration of this Act was limited to a period of ten years after its coming in force. In 1888, the Inebriates Act (51 and 52 Vict., ch. 19) was passed, which had for its object the re-enactment, with some slight modifications, of the Act of 1879. An amendment of the Habitual Drunkards Act of 1879 was passed in 1898 (61 and 62 Vict., ch. 60).

By these Acts "Retreats" are licensed for habitual drunkards with provision for medical attendance. If the person holding the licence is a duly registered medical practitioner, he may himself act as medical attendant to the Retreat. No such licence is given to any person who is licensed to keep a house for the reception of lunatics.

An habitual drunkard is defined by the Act as —

“ A person who, not being amenable to any jurisdiction in Lunacy, is notwithstanding, by reason of habitual intemperate drinking of intoxicating liquors, at times dangerous to himself or herself or to others, or incapable of managing himself or herself, and his or her affairs ”

Any habitual drunkard desirous of being admitted into a retreat may make a written application for admission, couched in the terms of the schedule appended to the Act of 1879, to the licensee of a retreat. The time the applicant undertakes to remain in such retreat must be specified in the application, which must be accompanied by a statutory declaration of two persons, stating that the applicant is an habitual drunkard. The applicant's signature must be attested by a justice of the peace, who before attesting must satisfy himself that the applicant is an habitual drunkard within the meaning of the Act, and must explain to him the effect of his application, and state in writing that he understood such explanation. The applicant after admission into a retreat will not be allowed to leave it till the expiration of the time mentioned in the application, provided that it shall not exceed a period of two years. At any time after the reception of an habitual drunkard into a retreat he may, upon the request in writing of the licensee of the retreat, be discharged by order of a justice, if the request is deemed by him to be reasonable and proper.

A person who is or has at any time been detained in a retreat may have his term of detention extended, or be re-admitted, in like manner as an habitual drunkard may be admitted, except that the statutory declaration (mentioned in the Act of 1879) shall not be necessary, and that the attesting justice shall not be required to satisfy himself that the applicant is an habitual drunkard.

At the request of the licensee the habitual drunkard at any time after his admission may, by permission of a justice, be allowed to live with any trustworthy person who is willing to take charge of him for a definite time, not exceeding two months, the leave may be prolonged every two months for a like interval until the whole period of detention is expired. Any time thus spent away from the retreat is deemed part of the time of detention, except where the leave of absence is forfeited or revoked.

If an habitual drunkard while detained in a retreat wilfully refuses to conform to the rules, he is liable on summary conviction to a penalty not exceeding five pounds, or, at the discretion of the Court, to be imprisoned for a period not exceeding seven days, and then returned to the retreat, the period of imprisonment being excluded from the original time he agreed to place himself under restraint. If he escapes he may be apprehended and remitted to the retreat, the time between his escape and return shall not be treated as part of his term of detention.

These Acts apply, with certain provisions as to technical procedure, to Scotland and Ireland, as well as to England.

So far as restraint goes, the **distinction** between a **lunatic** and a non-criminal **habitual drunkard** consists in the fact that the lunatic may be placed under restraint against his will, the habitual drunkard can only be detained with his own consent and for a limited time. It is necessary that an habitual drunkard should be sober at the time he makes the application before the justices.

**Criminal Drunkards.** Special provisions applying to habitual drunkards convicted of offences are contained in the Amending Act of 1898, of which the following are the more important —

" 1 (1) Where a person is convicted on indictment of an offence punishable with imprisonment or penal servitude, if the court is satisfied from the evidence that the offence was committed under the influence of drink or that drunkenness was a contributing cause of the offence, and the offender admits that he is, or is found by the jury to be, an habitual drunkard, the court may, in addition to or in substitution for any other sentence, order that he be detained for a term not exceeding three years in any State inebriate reformatory or in any certified inebriate reformatory, the managers of which are willing to receive him "

" 2 (1) Any person who commits any of the offences mentioned in the First Schedule to this Act, and who within the twelve months preceding the date of the commission of the offence has been convicted summarily at least three times of any offences so mentioned, and who is an habitual drunkard, shall be liable upon conviction, or indictment, or if he consents to be dealt with summarily on summary conviction, to be detained for a term not exceeding three years in any certified inebriate reformatory the managers of which are willing to receive him "

**Delirium tremens** being legally regarded as a diseased condition of the mind may render the subject of it a fit and proper person to be summarily placed under restraint. As a rule, the attack is too brief to necessitate the patient being sent under urgency order and certificate to an asylum, it is generally sufficient if he is restrained in his own house. In either case the position of the medical practitioner is uncertain as regards the steps that possibly may be taken by the patient after recovery, actions for damages have been brought against medical practitioners for having improperly deprived the plaintiff of his liberty and for ill-treatment whilst thus detained.

In one such case, *Scott v Wakem* (Guildford Ass, 1862), the plaintiff, who undoubtedly had been dangerous to others during his attack, and in consequence was taken charge of by a man placed in the house by order of the doctor, brought an action of this kind. The judge, Bramwell, B., ruled that if the plaintiff at the time was a dangerous lunatic and likely to do mischief to any one, the defendant would be justified in putting restraint upon him until there was reasonable ground for believing that the danger was over. Or, if the wife of the plaintiff had called in the defendant to cure her husband from delirium tremens, and that he did so, and left him when he believed he had recovered, he would be justified in what he had done provided he had done nothing that was not necessary nor reasonably proper under the circumstances. Notwithstanding this summing up a verdict was given against the defendant with, it is true, only a farthing damages, but the expense and annoyance of such a trial is no slight infliction. In another case, *Symn v Fraser and Andrews* (Queen's Bench, 1863), a similar action was brought a year after the alleged ill treatment and restraint was put in force. In this case the verdict was for the defendant. Cockburn, C. J., before whom the case was tried, said, that even if attendants or nurses are not originally appointed by a medical man, yet if he assumes authority and command over them in reference to the management of the patient, he would be responsible for the personal restraint under which the patient was placed.

From this it will be seen that the position of the medical practitioner in charge of a case of delirium tremens is by no means secure, if he neglects to take proper precautions, and harm ensues, he is blameworthy, and if he takes these precautions he is liable to be proceeded against. If the medical attendant in a case of delirium tremens performs his duty to his patient in a proper manner he need not fear an adverse verdict should an action for damages be brought against him, he cannot, however, avoid the possibility of such an action being brought. If he is dealing with people who are strangers to him, or he sees any probability of subsequent trouble, it is well to have an authorisation in writing from the husband, wife, or other near relation, to do what is necessary as to

restraint Such an authorisation will not guarantee the medical attendant against an action, but it will be of material value in defending it In the case of *Scott v Wakem*, already quoted, the wife denied having authorised the defendant to interfere, although when he was called in he found the plaintiff with loaded pistols in his hands threatening to shoot her, in this case the facts contradicted the wife's statement, but it might happen otherwise

If a patient with delirium tremens is duly certified, and in consequence has been placed under restraint, an action entered against the medical practitioner who gave the certificate would probably be stayed upon summary application to the High Court, or a judge thereof, as provided in the Lunacy Act, 1890, but the medical practitioner would have to satisfy the Court that there was no reasonable ground for alleging want of good faith or reasonable care



## PART III.—TOXICOLOGY

## CHAPTER XXVIII

## POISONS IN THEIR GENERAL ASPECT.

TOXICOLOGY is that branch of science which relates to poisons, it is generally understood to include a description of the symptoms and treatment of poisoning, as well as of the nature, chemical constitution, and methods used in the isolation and detection of poisons. An accurate and, at the same time, terse definition of the word poison is not readily attained. In the broad sense, a poison is a substance which on being absorbed into the living organism, or by its chemical action on the tissues, injures the health, or destroys life. Such a definition would include pathological ferments, which, though regarded as poisons from the medical standpoint, do not come within the range of toxicology. Again, many substances when absorbed into the organism in large quantity are deleterious to health, and yet they can hardly be regarded as poisons, since the amount in which they solely become injurious is so excessive that they may be taken with impunity in much larger doses than is the case with a substance which is recognised as a poison. Any substance that is not absolutely inert would produce injurious effects if administered in excess, so that noxious activity has to be taken into account. A substance which produces no ill effects when given in a single small dose, may do so if similar doses are repeatedly administered. This is a question of practical moment to the toxicologist in relation to the frequent reception by the organism of minute quantities of poisons, such as arsenic, lead and mercury, the amount introduced on any one occasion may be insufficient to produce an injurious result, but if taken day after day the repeated small doses would eventually be harmful. Some substances, such as the mineral acids, when swallowed destroy tissues without being absorbed, and in this way may cause death by "poison."

In cases of criminal poisoning it is no longer necessary to determine whether the substance administered comes under the definition of a poison, the law relating to this subject (24 and 25 Vict., ch. 100, section 11) runs thus —

"Whosoever shall administer, or cause to be administered or taken by any person, any poison or other destructive thing, with intent to commit murder, shall be guilty of felony."

Two points demand attention that the intent to commit murder determines the culpability of the act, and that the comprehensive words, "or other destructive thing," makes it immaterial whether the substance given comes under the usual definition of a poison or not. The question of amount has to be considered. Lord Chief-Justice Cockburn ruled in *Reg. v. Hennah* (Cornwall Assizes, 1877), that unless the thing administered was in sufficient amount to be noxious, it does not come under the legal definition of a noxious thing,

and, further, that a distinction is to be drawn between a thing that is only noxious when given in excess, and one that is recognised as a poison and known to be noxious and pernicious in its effect. This ruling throws great responsibility on the medical witness, for, although he may not be called upon to define a poison in the abstract, he will be required to express an opinion as to whether the substance given was a noxious substance and whether it was given in excess. The law is very comprehensive in dealing with the culpability of an intentional poisoner, thus 21 and 25 Vict., ch. 100, s. 23, states:—

“Whosoever shall unlawfully and maliciously administer to, or cause to be administered to, or taken by any other person, any poison or other destructive or noxious thing, so as thereby to endanger the life of such person, or so as thereby to inflict upon such person any grievous bodily harm, shall be guilty of felony.”

The amount of damage sustained that would constitute “grievous bodily harm” is capable of being variously estimated and the medical witness may have great difficulty in conveying his opinion to the jury through the media of examination and cross-examination only, as to whether the substance, in the dose in which it was given, could or could not cause such harm. In respect to the administration of many substances a variety of conditions determine the intensity of the noxious effects which may ensue. When the prosecutor has not received any harm from the alleged “noxious thing,” the medical witness may be asked questions with regard to hypothetical cases of which a full description of the conditions and surroundings is not afforded, opinions given under such circumstances are liable to mislead the jury.

The law provides that the administration of any poison, or other destructive or noxious thing, with the *intent* to injure, aggrieve, or annoy constitutes a misdemeanour, and, further, that if a prisoner is charged with felony for having administered a poison or other noxious thing, and the jury are not satisfied that he is guilty of felony, but are satisfied that he is guilty of a misdemeanour as above, they may find him guilty of the lesser crime.

In cases of poisoning, therefore, medical evidence is not limited to the bare statement that the substance administered is possessed of poisonous or noxious properties, it must go further and establish the dose at which the substance becomes poisonous or noxious. Several conditions require to be considered in determining the toxic effects of poisons, among which are *age*, *idiosyncrasy*, *habit* and *state of health*.

**Age.** Young children succumb more easily to the effects of poison than adults, this is especially the case with morphine. On the other hand, children tolerate a poison like belladonna better than adults, young adults tolerate the action of irritants, such as tartar emetic, better than old people.

**Idiosyncrasy.**—Some persons are naturally tolerant, and others intolerant, of various substances which have poisonous properties. In some cases an ordinary full medicinal dose of arsenic acts as a poisonous dose, and causes gastro-intestinal irritation. Medicinal doses of strychnine occasionally produce muscular twitchings, and even slight general spasm. Small doses of mercury salivate some people, whereas others can take it for a prolonged period without any apparent effect being produced. In some persons certain articles of food invariably produce symptoms of poisoning—nausea, vomiting, and diarrhoea—whilst in others the disagreeable effects are limited to an attack of nettle-rash, the majority of people being able to eat such food without any ill effects whatever.

**Habit.**—The regular and prolonged use of some poisons in small doses tends to develop tolerance on the part of the system to their action. Morphine,

tobacco, and alcohol are examples of poisons towards which tolerance may be thus acquired. Persons who in this way are habituated to the use of such substances can with impunity take what would be a poisonous dose to a novice.

**The State of Health.** In a broad sense, healthy individuals are less likely to succumb to the effect of poison than those who are weakened by disease; there are certain notable exceptions, however. Patients suffering from acute mania, delirium tremens, tetanus, or dysentery can take an amount of opium that might be sufficient to cause the death of a non-habituated healthy person. On the other hand, opium is exceedingly dangerous in granular kidney, apoplexy, and hyperæmic diseases of the lungs. Digitalis, tobacco, and tartar emetic are badly tolerated in cases of fatty or weak heart. Any irritant would aggravate the symptoms of gastro-intestinal catarrh. Gouty subjects, especially if suffering from granular kidney, are intolerant of repeated small doses of lead.

The **physical condition** of the poison and the **mode** of its **introduction** into the system exercise a considerable influence on the rapidity and the intensity of its action. Poisons that are capable of assuming the **gaseous** form act much more immediately and energetically when thus administered than when given as **liquids** or **solids**. Arsenic, if inhaled in the gaseous form as arsenuretted hydrogen, acts more rapidly than when arsenious acid dissolved in water is swallowed, solid arsenious acid is still less active. The **mode of administration** largely influences the effects produced by a given dose of the poison; its physical condition remaining the same. A solution containing a poison acts most promptly if injected directly into the blood current, less rapidly if brought in contact with the serous membranes, cellular tissue, mucous membranes, and the skin; the rapidity of absorption diminishes in the order of the tissues named, absorption from the intact skin being the slowest of all.

Two points have to be considered in relation to the effect produced by a given dose of a poison, the rapidity of **absorption** and the rapidity of **elimination**. If a poison is eliminated as rapidly as it is absorbed, and the rate of absorption is too slow to immediately bring the system under its lethal influence, no permanently deleterious effects are produced. For this reason some poisons can be received into the mouth and even swallowed with impunity, in amounts that would be lethal if introduced under the skin; curare, some kinds of arrow-poison, and the venom ejected from the fangs of poisonous serpents are examples of poisons of this type. The wound inflicted by a poisoned arrow may have a sufficient amount of the poison deposited in it as to cause the death of the wounded person if left to his fate, but if a second person immediately and vigorously sucks the wound, no mischief results to either of them. The same immunity follows if the wounded individual himself is able to suck his wound, and does so at once; the poison extracted is at once spat out, and thus only momentarily comes in contact with the mucous membrane of the mouth, the absorptive capacity of which is too limited to allow of the introduction into the system of an amount sufficient to cause mischief. The poison extracted by the suction might probably be swallowed with impunity, as it would be eliminated by the kidneys as quickly as it would be absorbed by the gastric mucous membrane.

The law makes no distinction in relation to the mode in which a poison is introduced into, or acts upon, the system. Hypodermic, or endermic administration of a poison, or the injection of it into the bowel, with intent to commit murder, are regarded precisely as though the poison had been given by the mouth.

The **chemical combination** in which certain poisonous substances exist, and the degree of **concentration** in others, exercise a powerful influence on their lethal potency. Silver nitrate and hydrochloric acid are both energetic poisons when taken separately, but when combined the resulting salt silver chloride is inert, or thereabouts, because of its insolubility. In some instances the action of the poison is entirely altered by chemical combination, another effect being substituted for that which is characteristic of it in the uncombined state. Strychnine alters the rate of transmission of stimuli from cell to cell in the spinal cord, or possibly increases the excitability of the cells, the result being the occurrence of clonic muscular spasm. If strychnine is so treated as to become a methyl derivative, it no longer acts on the cord, but it paralyzes the motor nerve endings, like curare. In the case of poisons, such as the mineral acids, which act directly on the tissues with which they come in contact, the degree of concentration is an important factor. An amount of a concentrated mineral acid that would be sufficient to cause death, might be swallowed with impunity if largely diluted with water.

When poisons are taken by the mouth, the state of the stomach as regards **presence or absence of food** considerably modifies the rate of absorption and the intensity of local action, a person has been known to swallow more than a lethal dose of undissolved white arsenic with impunity because the stomach was well filled with oatmeal porridge. When a poison is received into a full stomach the usual symptoms are delayed much beyond the customary time of onset, and if the poison is one that is quickly eliminated by the kidneys an average fatal dose may be survived although the whole of it is eventually absorbed, on the other hand, an empty stomach and bowel quickly absorb any poison received, and the onset of the symptoms is accelerated.

**Classification of Poisons.** A comprehensive classification is scarcely feasible without involving great complexity and, when accomplished, is of little use. The division of poisons into **inorganic** and **organic**, with the subdivision of the first into *corrosive* and *irritants*, and of the second into *irritants* and *neurotics*, affords a basis sufficiently broad for practical purposes. Both organic and inorganic groups comprise poisons which act by disintegrating, or by interfering with the functions of the red corpuscles of the blood.

### THE DIAGNOSIS OF POISONING.

The effects severally produced on the living organism by various poisons differ within limits in accordance with the kind of poison taken. Common to almost all forms of poisoning, however, are a number of symptoms, or effects, some of which are intrinsic, affecting the individual, others are accidental and do not personally implicate him.

In the first group is **sudden occurrence** of acute symptoms in a person previously in his usual health. In relation to this indication it is to be remembered that the initial symptoms produced by many poisons bear a certain **resemblance** to those due to **disease**. Arsenic causes symptoms which have been mistaken for those of cholera or of gastro-intestinal catarrh, strychnine for those of tetanus, morphine for those of apoplexy, belladonna for those of acute delirious mania or of alcoholism. Errors of the opposite kind have been fallen into the rapid onset of an **acute disease** in a person who appeared to be in good health up to the time when the symptoms showed themselves has been **mistaken for the effect of poison**. Among diseases of this nature are—an acute gastric ulcer which ruptures, perforation of the bowel, rupture of an abdominal aneurism,

the formation of a peri-uterine hæmatocele, acute intestinal obstruction with persistent vomiting, choleraic diarrhœa hæmorrhage into the pons. Some of these conditions could only give rise to momentary doubt, others may cause prolonged anxiety to the medical attendant.

The occurrence of symptoms of an anomalous nature **shortly after eating** or drinking or taking medicine is a suspicious indication, which for its interpretation requires considerable discrimination. Violent vomiting and purging may be due to some change undergone by the food itself before cooking. The food may be in its normal condition, but the state of the recipient's stomach may be at fault, a person who has fasted long beyond his usual meal time, if he hurriedly eats some food difficult of digestion, may forthwith be attacked by vomiting and pain which closely resemble the effects of an irritant poison, an unsuspected gastric ulcer may give way under like conditions. Poison may be intentionally added to medicine which is being taken by a sick person, and the ill results which follow every dose may be attributed by him to its legitimate therapeutic action. When attending a case in which anomalous symptoms repeatedly recur, the medical practitioner should be silently observant, unless the indications of foul play are conclusive, or the state of the patient is becoming critical, in either of these events it is his duty at once to take such measures as are necessary to prevent further mischief. The position is an extremely delicate one. It is a very serious matter for a practitioner to make an unfounded charge against an innocent person, but to allow a patient to be poisoned under his eyes is infinitely worse. If the medical attendant's suspicions are aroused, he should especially distrust the ministrations of any one who is studiously attentive to the invalid, who prepares all his food and insists on giving it to him with his or her own hand, and who displays an exaggerated interest in the treatment of the patient and in the visits of the doctor. Such a person will stand by the invalid as he takes his food, and will throw away what is left, under the pretext that everything partaken of should be freshly prepared. All this is not inconsistent with innocence and a genuine desire for the welfare of the patient, but when incongruous symptoms arise which are at variance with the natural course of disease, such a person is to be regarded with suspicion and should be carefully watched. The patient should be cautiously, but not mysteriously, interrogated as to the food or fluids that he has partaken of, and the times when he felt any accession of the suspicious symptoms. In cases of suspected foul play the engagement of a couple of trained nurses for night and day duty respectively, affords the most practical safeguard, they should be instructed to prepare all food, to give it and the medicine with their own hands, and not to leave the patient when on duty. In giving strict injunctions to this effect, it is not necessary that the nurses should be taken into the confidence of the doctor, unless the condition of matters is critical, or for some other reason he deems it advisable. A twenty-four hours' supply of the patient's urine should be obtained and submitted to chemical examination for the suspected poison, this may be done without giving rise to suspicion, as doctors frequently require specimens of urine for examination in the course of ordinary diseases. To take away food or beverages prepared for the patient would, of course, be to reveal suspicion, and should only be resorted to when the conviction of foul play is very strong and concealment is no longer possible. If the urine, or other substance, is sent to an analyst, it should be accompanied by a statement as to the kind of poison it is suspected to contain.

The state of the medical attendant's mind when he believes that an attempt

is being made to poison his patient, may be described as having two stages first, one of suspicion, and then one of conviction, supposing that his suspicions are well founded. These states of mind have different obligations. When, from symptoms irreconcilable with the ordinary course of the disease, or on account of some suspicious circumstance or chain of events, the idea dawns upon the medical attendant that his patient is the victim of a secret poisoner, he is not therefore, justified in immediately proclaiming his suspicion, it is quite possible that he may be wrong, and as a statement to the effect that he believes some one is attempting to commit murder cannot be made without implicating people who may be innocent, it should not be rashly uttered. Besides doing an injustice, the practitioner would render himself liable to legal proceedings on the part of those whom he had directly, or by implication, accused of the crime. Errors of interpretation, or of judgment, are easily fallen into, and under circumstances such as are being discussed it would be unwarrantable, on the first feeling of mistrust, to act as though the matter was beyond all doubt. The steps to be taken when suspicion is aroused but no evidence has been obtained, are to protect the patient in the way previously described, and to keep a sharp look out for renewed attempts.

If the matter has gone further, and the medical attendant is fully convinced that poison is being administered, what is he to do? There are three courses open: to tell some member of the family other than the suspected person, to tell the patient himself, or to inform the police. A fourth plan has been recommended: it is to tell the person believed to be the culprit, that proof of the administration of poison has been obtained (without accusing him of being the administrator) and that, in the event of any further attempts being made, it will be necessary to inform the police; this would probably put an end to the matter, but it comes too near compounding a felony to be justifiable. Strictly speaking, it would be the duty of the medical attendant to inform the police as soon as he is fully convinced that a criminal attempt is being made on the life of his patient, but before doing so circumstances may make it advisable for him to take into his confidence a member of the family whom he can trust. Just as precipitancy is to be deprecated in the stage of mere suspicion so is promptness demanded when that suspicion is converted into certainty. The medical attendant's duty is clearly defined, and he is bound to fulfil it without fear or favour.

Some striking examples of the difficulty of distinguishing between disease and the effects of an irritant poison frequently administered in small doses were brought to light during the trial of Severino Klosowski, in 1903, for the murder of Maud Marsh, with whom he had lived. The deceased was taken ill in October, 1902, and was first seen by a medical man on the 10th of that month. She was found suffering from diarrhoea, vomiting, and great pain in the stomach, and a provisional diagnosis of gastro enteritis was made. Two days later she was very much better, but on the third day the symptoms recurred with great severity and there was rigidity of the muscles of the leg. The patient became steadily weaker, and on October 20 a consultation was held with another medical man, when an opinion was formed that she was suffering from irritant poisoning, probably ptomaine. After they had separated one of the medical men became suspicious of arsenical poisoning, but it was then too late to save the woman, who died the following day. An inquest was held, and at the post mortem antimony was found in the contents of the stomach and bowels, and in the liver and kidneys, the total amount of tartar emetic in the whole body being estimated at from twenty five to thirty grains. It was also shown that Maud Marsh had been admitted into Guy's Hospital in July, 1902, with symptoms of vomiting, pain and diarrhoea, and a diagnosis of peritonitis had been made, but no cause found for it.

The bodies of two other women with whom the prisoner had lived were then exhumed, and an analysis made of the viscera and their contents.

The first was that of Mary Chapman, who died on December 25, 1897, and was exhumed

on December 9, 1902. The body was found to be remarkably well preserved, "the face and head were those of a woman who might have been confined that day." The spleen, the kidneys, the bladder and the heart were all normal. The stomach and bowel showed signs of gastro enteritis. A large amount of antimony was found in the liver, kidneys, stomach, and intestines. The death had been certified as due to phthisis.

The second woman, Bessie Taylor, died on February 13, 1901, and was exhumed on November 22, 1902. The body showed no putrefaction and no odour. The organs were all normal except the stomach, where indications of gastritis were present. The inner surface of the bowel was coated with the yellow sulphide of antimony. There was no affection of the uterus or signs of intestinal obstruction. Some thirty grains of tartar emetic were recovered from the stomach, liver, and kidneys. In this case medical evidence relating to the illness of the deceased was given. She was first seen by a medical man (the same who later attended Maud Marsh) on January 1, 1901, and was suffering from diarrhoea and pains in the stomach. Remarkable variations in the course of the illness occurred, the patient being much better on some days, and unexpectedly relapsing on others. One day she was up playing the piano, and the next was as bad as ever. Three separate consultations with three other doctors were held. One of these, a specialist in diseases of women, suggested that the patient was suffering from an affection of the uterus, another thought the symptoms were due to a severe form of hysteria, and the third diagnosed cancer of the stomach or intestines. Death occurred on February 13, and the cause was given in the certificate as intestinal obstruction, vomiting, and exhaustion.

Klosowski was found guilty and was executed.

**Indications of poisoning** may be shown by circumstances **apart from** the symptoms which affect **the individual**. A number of healthy persons may be simultaneously attacked with analogous symptoms after a common meal. This may be due to intentional or to accidental admixture of poison with the food, or to abnormal constitution of one or more of the food-stuffs partaken of, suspicion may also be aroused by the discovery of an unusual appearance or odour of the patient's food or medicine.

When a medical man is in attendance on a case in which he suspects poisoning, he should note down all that he observes immediately after each visit, such notes may be of the utmost value should the case end fatally. If the conviction of foul play is believed to be well founded, medicine bottles, specimens of foods and drinks provided for the patient should be taken possession of, and guarded until handed over to the police or to the analyst, vomited matter, and sheets or other fabrics stained therewith, should also be unpounded.

## THE GENERAL TREATMENT OF POISONING.

The objects aimed at are to remove from the digestive tract any poison it may contain, or to neutralise it, to combat the effects of that which has been absorbed, and to promote its elimination, to keep the patient alive until the effects of the poison have passed off, to alleviate general symptoms.

There are two ways of forcibly emptying the stomach—by means of an *emetic*, and by means of the *stomach-tube*.

**Emetics.**—Half-drachm doses of zinc sulphate dissolved in warm water repeated if necessary, act quickly without causing depression, if not at hand, a dessert-spoonful of mustard may be given in a tumbler of warm water. For children, a teaspoonful of ipecacuanha wine is a good emetic. In any case the patient should drink copiously of warm water, which materially aids emesis and washes out the stomach. Tartar emetic and copper sulphate are to be avoided, the former is a depressant, and both would tend to add to the difficulty of a subsequent chemical analysis, should it be necessary, and exception is to be made with regard to phosphorus poisoning, in which copper sulphate may be given. Instead of giving an emetic by the mouth, a hypodermic injection

of a solution of apomorphine hydrochlorate may be administered, the B P solution for hypodermic injection contains one grain in one hundred and ten minims, of which ten minims one-tenth of a grain is a proper dose to inject in a case of poisoning. This method of procuring emesis is convenient, especially in cases of narcotic poisoning where there is great difficulty in making the patient swallow. In the absence of an emetic, the fauces may be tickled with a feather, or even with the finger, and copious draughts of warm water given.

The **stomach-tube** is an efficacious instrument for emptying the stomach independently of physiological action, and is, therefore, entirely under the control of the operator. If the mouth of the patient can be kept open, there is little difficulty in passing the tube of the instrument down the œsophagus, keeping it well against the posterior wall, it may be necessary to open the jaws forcibly, and to keep them open with a gag. Before withdrawing any of the contents of the stomach, a pint or so of warm water should be injected, the same amount is then withdrawn, and a further supply injected, which is also withdrawn, the object being to wash the stomach out without entirely emptying it, which might lead to injury of its coats. If the stomach-tube is not to hand, five or six feet of india-rubbing tubing such as is used for small gas supplies, fitted with a funnel at one end, may be substituted. The free end is passed down the patient's throat, and, when in the stomach, a pint or more of warm water is poured into the funnel, which is held above the level of the patient's mouth. When the funnel is almost empty the tube close to it is pinched between the finger and thumb, and the funnel depressed until it is lower than the stomach, on removing the finger and thumb the tube acts as a syphon, and evacuates the stomach. The process is to be repeated as with the stomach-tube until nothing but clear, non-odorous water comes away. If the stomach contains much solid matter of a lumpy consistence, it may be advisable to give an emetic before using the pump or tube, in order to avoid clogging. *Neither the stomach-tube nor emetics are to be used in cases of corrosive poisoning*; the proper treatment in such cases is to neutralise the poison, carbolic acid, however, constitutes an exception to this rule. The passage of the tube is exceedingly risky if the walls of the œsophagus and stomach are softened and corroded, for this reason, the stomach-tube requires using with great caution in the case of certain irritants, especially if the patient is not seen for some time after the poison is swallowed. In cases where the proper treatment is to empty the stomach, and the introduction of the stomach-tube would be risky, an emetic should be given. The stomach-tube is especially useful in cases of poisoning by opium, alcohol, chloral hydrate, chloroform in the liquid state, the vegetable, and most of the mineral irritants, phosphorus (if the case is seen shortly after the poison is swallowed), and the alkaloids. In strychnine poisoning it will probably be necessary to place the patient under chloroform before the tube can be passed. In the absence of the stomach-tube an emetic may be administered in appropriate cases, but where time is of importance the mechanical method is preferable.

If spontaneous purging has not occurred it may be necessary to evacuate the bowels by means of purgatives or by enema.

**Antidotes** are remedies which counteract the effects of poisons. They act either *mechanically, chemically, or physiologically*.

Flour and water, and chalk mixture, act as **mechanical** antidotes, when given in poisoning by phosphorus or cantharides. Magnesia and chalk are **chemical** antidotes to the mineral acids, as are the alkaline sulphates to the salts of lead and barium. **Physiological** antidotes act upon the tissues or organs



Jona has shown by experiments on animals that the administration of adrenalin delays the absorption of certain poisons, for example, cyanides, strychnine, and aconite, by its vaso-constrictor action upon the gastric mucosa<sup>1</sup>. The subject is further considered under the heading of "Antagonism of Poisons."

The **elimination** of the poison that has been absorbed is to be assisted by purges (when not contra-indicated), diuretics, and special remedies in the case of certain poisons.

Every endeavour must be made to keep the patient alive until the effect of the poison has passed off by artificial respiration and cold douche (hydrocyanic acid), by being kept awake (opium), by external warmth (chloral hydrate and carbolic acid), and by stimulants.

**General symptoms**, as excessive pain, exhaustion, useless vomiting, and purging, are to be combated by appropriate remedies.

### ANTAGONISM OF POISONS.

This term is applied to the power certain poisons are supposed to possess of counteracting the effects of other poisons, either by the direct exercise of an opposing influence—if a poison paralyses a certain tissue, its antagonist stimulates it—or, as suggested by Ringer,<sup>2</sup> by chemical displacement. The latter hypothesis supposes that a poison which acts as an antagonist to another poison has a stronger affinity for the tissue attacked, and that it displaces the poison towards which it is an antagonist, substituting its own action for that of the poison.

In its full meaning the term "antagonism" includes more than mere reversal of some of the effects produced by a poison, it comprises counteraction of the influence of the poison step by step in the tissues originally attacked. This is altogether different from setting up at a distance an opposing force or obstacle, which only changes or reverses outward indications, leaving the tissues originally attacked still under the influence of the primary poison—merely blocking the way to external manifestations. For example, morphine slows the action of the heart and atropine quickens it, therefore, as far as outward appearances go, atropine in this respect acts antagonistically to morphine. But morphine slows the heart by excitation of the vagus at its origin in the brain—as shown by the fact that if the vagi are divided before morphine is given, no retardation occurs. Atropine quickens the heart's action by paralysing the terminations of the vagi, and also inhibitory ganglia in the heart—as shown by absence of slowing on irritation of the vagi in animals under the influence of atropine. Under these circumstances it is clear that the influence exerted by morphine, in the direction of slowing the heart, is not removed by atropine, it is simply arrested at a point distant from the seat of action.

Again, a true antagonist would counteract in every direction the influence of the poison to which it is opposed, it is not enough for it to combat some effects, and to leave others unopposed. Atropine appears to some extent to be antagonistic to morphine as regards the respiratory function: atropine stimulates the respiratory centre for a time and then like morphine depresses it. Unverricht,<sup>3</sup> however, denies that atropine is a stimulant for the normal respiratory apparatus, and cites some experiments by Orłowski<sup>4</sup> in support of his views. Atropine also at first stimulates but subsequently depresses the

<sup>1</sup> *Brit Med Journ*, 1913.

<sup>2</sup> *Handbook of Therapeutics*, 1888.

<sup>3</sup> *Centralbl f klin Med*, 1891 and 1892. *Berliner klin Wochenschr*, 1896.

<sup>4</sup> *Einwirkung des Atropins auf die Resp*, 1891.

vaso-motor centre, opium in large doses depresses it from the first. In regard to other functions the action of the two poisons, though apparently antagonistic, is not really so, and in others again there is not only absence of antagonism, real or apparent, but similar results are produced, though not in the same manner. As is well known, opium contracts the pupil and atropine dilates it, opium acting centrally and atropine peripherally—paralysing the terminals of the oculo motor nerves. Both poisons cause dryness of the mouth—opium by lowering reflex excitability, and atropine by paralysing the secretory fibres of the chorda tympani. Opium causes sweating by stimulating the central nerve-apparatus concerned, atropine arrests it by paralysing the nerve terminals in the sweat glands. Opium, after causing an initial increase in the intestinal movements, arrests them probably by lowering the reflex excitability, atropine is believed ultimately to deprive the intestines of movement by paralysing their motor nerves and, finally, the muscular elements themselves, the terminals of the inhibitory fibres of the splanchnics are also paralysed by atropine.

Thus, although in respect to one or two prominent symptoms, atropine and morphine are opposed in their action, they are not true antagonists. Their modes of action are different, broadly speaking, *atropine acts peripherally, morphine centrally*. Morphine, in poisonous doses, depresses the excitability of the ganglionic cells of the cerebrum, and probably of the cord also and lessens reflex function. This is disputed by some, Unverricht<sup>1</sup> states that opium does not lower, and may increase, the irritability of the cortex. Atropine stimulates the central nervous system, and in this way increases the reflex function, but it also paralyses many of the peripheral nerves, and thus cuts off the organs supplied by them from their centres. The ultimate effect of both poisons is to paralyse the motor and the sensory nerves. Morphine, when used as an antagonist to atropine, is much feebler in its effects than atropine is when used to antagonise morphine.

Notwithstanding the absence of proof of antagonism, many cases are recorded in which life is believed to have been saved by the administration of atropine to patients suffering from poisonous doses of opium, but a careful study of these cases occasions great doubt as to the correctness of the inference drawn by those who advocate the treatment. When active treatment is successful, there is a tendency to attribute the good results to some specific cause, and the remedy, or the means used, that is apparently the most active gets all the credit. It is quite excusable for a medical man, after administering atropine, to be greatly impressed by the recovery of a patient from an apparently moribund condition, the result of a poisonous dose of opium. It is not customary, however, in such cases to trust to one remedy solely, no matter how much importance is attached to it, other treatment is actively carried out during the critical period, and the effects of such treatment are often ignored, or underrated. There are grave reasons for believing that too active treatment of opium poisoning with atropine has sometimes been the means of hastening, rather than preventing, the fatal issue. Lenhartz<sup>2</sup> states that out of 132 cases of opium poisoning, of which 59 were treated with atropine, and 72 without, 38 per cent of the former died, and only 15 per cent of the latter. In three cases that came under his own observation, subcutaneous injection of atropine produced no effect beyond dilating the pupils and increasing the rapidity of the heart beats, the unfavourable symptoms persisted, recovery ultimately took place in two, the third died, and the cumulative effect of the atropine was regarded as not

<sup>1</sup> *Centralbl. f. klin. Med.*, 1891

<sup>2</sup> *Archiv f. exp. Path. u. Pharm.*, 1887

being beyond suspicion of having exercised a deleterious influence. Bashford's<sup>1</sup> experiments on animals show that when the antagonising dose of atropine is above one thirty-sixth of its minimum lethal dose it is useless and often hastens death by adding the lethal effects of atropine to those of morphine. He recommends that not more than one-twelfth of a grain of atropine should be injected, and that the dose should not be repeated. The earlier the atropine is administered the greater is its effect. External warmth materially aids the antagonistic action of atropine in morphine poisoning.

Without going so far as to deny the possibility of benefit from the use of atropine in the treatment of opium poisoning the exercise of great caution is strongly urged, as the repeated heroic doses that have been given are altogether unjustifiable in the face of the known ultimately combined effects of the two poisons, it is to be remembered that whatever antagonistic effect small doses of atropine may exercise, its ultimate effect is to paralyse the heart.

Perhaps the best illustration of antagonism between poisons is afforded by atropine and physostigmine. Fraser's<sup>2</sup> experiments have demonstrated that physostigmine increases the excitability of the vagi, while atropine diminishes and suspends it, physostigmine lowers arterial tension, atropine augments it, physostigmine increases glandular secretion, atropine diminishes or arrests it, physostigmine contracts the pupils, atropine dilates them. In the greater number of these instances the antagonism is real, being effected in the same structures and in the same order.

Limited antagonism between poisons may be advantageously utilised for antidotal purposes. The effects produced by strychnine on the respiratory centres and on the reflex mechanism of the cord are, within limits, capable of being antagonised by chloral hydrate. Reciprocally, chloral poisoning may be beneficially treated by the administration of strychnine, atropine is also a limited antagonist to chloral hydrate. Muscarine antagonises atropine by stimulating the endings of the oculo-motor nerves, the endings of the secretory fibres of the chorda tympani, and the inhibitory cardiac ganglia, at the same time paralysing the cardiac muscles, it also depresses the activity of the respiratory centres. Muscarine ultimately paralyses the inhibitory action of the vagi, and is thus finally in accord with, instead of antagonistic to, atropine.

Other poisons are accredited with a limited degree of mutual antagonism, such as atropine to aconite, digitalis to aconite, chloral hydrate to picrotoxine, atropine to pilocarpine, aconite to strychnine, morphine to hyoscyamine, strychnine to nicotine. Theine and its congeners, caffeine and guaranine, to some extent antagonise morphine.

## GENERAL SYMPTOMS OF CORROSIVE AND OF IRRITANT POISONING.

Before considering the evidences of poisoning that are available in the dead body, it will conduce to their due appreciation if a general description is first given of the symptoms which occur during life. The poisons which give rise to the most characteristic post-mortem appearances are **corrosives** and **irritants**.

A **corrosive**, as the name indicates, is a substance which destroys tissue by direct chemical action, a corrosive also acts as an irritant, and, if administered in a dilute form, it may do so exclusively, without producing any corrosive

<sup>1</sup> *Archiv Internat de Pharmacodyn et de Therap*, 1901

<sup>2</sup> *Transactions of the Roy Soc Edin*, vols xxiv, xxvi

effects When a poison which acts as a corrosive is swallowed, an *immediate* and violent sensation of pain is produced, which extends from the mouth, along the œsophagus to the stomach, from whence it radiates over the abdomen. Uncontrollable retching and vomiting occur within a few minutes, the appearance of the vomited matter being determined within certain limits by the nature of the corrosive, shreds of mucous membrane, coagulated mucus, and blood are always present, the colour of the blood being sometimes altered by the chemical action of the poison. During this first stage the patient is frequently partially or wholly convulsed, a reflex symptom caused by the excruciating pain. There is intense thirst, with difficulty or impossibility of swallowing, each attempt to do so causing increased vomiting. The patient is in a condition of extreme collapse the surface, pale and cold, is bedewed with clammy sweat, the features are pinched, the eyes, sunk into their sockets, have a wild, terrified look. The voice is hoarse, or there may be complete aphonia, in the latter case it is probable that some of the corrosive has reached the larynx. The mouth is filled with ropy mucus, the salivary glands secrete profusely, the lips are swollen, and, along with the corners of the mouth, may show signs of the local action of the corrosive, the mucous membrane of the mouth is detached and the underlying tissues are corroded, the colour of the surface varying with the nature of the corrosive. The abdomen is usually distended. The breathing is laboured and noisy, attempts to clear the air-passage give rise to a distressing cough, which has a peculiarly hoarse, laryngeal sound. The pulse, thread-like and of low tension, is scarcely perceptible at the wrist. The bowels are confined, the urine is diminished in amount, or is entirely suppressed, and attempts to relieve the bladder are painful and futile. The mind usually remains clear to the end death taking place from extreme collapse, in some cases death is preceded by convulsions. In an acute case, as above described, death takes place **within twenty-four or thirty-six hours.**

An *irritant* poison is one which by its specific action sets up inflammation in the intestinal tract. A pure irritant does not produce corrosion, although some substances classed as irritants may act as corrosives. When a substance that acts solely as an irritant is swallowed, the symptoms do *not* come on in the act of swallowing *nor immediately after*, as is the case with corrosives, an interval elapses of from half an hour to an hour or more. In the case of metallic irritants (the symptoms they produce being taken as a type of irritant poisoning) there may, or may not, be an astringent or metallic taste perceived at the time the poison is swallowed, the presence or absence of this symptom depends partly upon the nature of the poison itself, and partly upon the medium in which it is administered. The first symptoms are those of gastro-intestinal irritation - violent and persistent vomiting and purging, with severe gastric and abdominal pains, the vomited matter probably at first consists of food, then it becomes bilious, and finally it may be blood-stained. There is intense thirst, and attempts to allay it provoke further vomiting. A hot burning sensation along with a feeling of constriction is felt in the throat. The purging is accompanied by violent tenesmus, and the dejections may be blood-stained, they are sometimes colourless, of the rice-water type. The symptoms of collapse appear the surface is cold and clammy, and the pulse feeble and intermittent, occasionally the skin is hot and dry, probably due to an attempt at reaction. There is great restlessness and anxiety, the mind often remaining clear to the last. The patient may be troubled with violent cramps in the legs, or he may have general convulsions. In fatal cases, death from exhaustion takes place **in from one to four days.**

**BLOOD POISONS.**

Many poisons exercise a remote influence on the blood, but by a blood poison is meant a substance which acts directly and specifically on the blood and through it affects the tissues at large. Some poisons have a composite action: they produce definite changes in the blood, but they also attack other tissues upon which possibly their chief toxic energy is expended, the pure blood poison attacks the blood only. The blood poisons may be divided into two classes: I Those which act primarily on other tissues besides the blood, II Those which for the most part act primarily on the blood only.

Much need not be said about Class I, the members of it not being recognised essentially as blood poisons. The concentrated mineral acids and strong solutions of the alkalies belong to this class. When strong sulphuric acid is swallowed it has a limited and local action on the blood, converting the hæmoglobin within the blood-vessels of the stomach into hæmatin which blocks the lumen of the vessels, in some instances this conversion into hæmatin has been observed to extend to the vena cava even as far as the right side of the heart. Poisonous doses of the mineral acids act as blood poisons in another way: they reduce the alkalinity of the blood and thus interfere with its power of carrying  $\text{CO}_2$ . Some of the heavy metals also belong to this class, as, for example, lead and mercury, which however, only act as blood poisons when taken in repeated doses for a considerable time.

Class II includes all the typical blood poisons which act on the entire mass of blood, and for the most part primarily on it alone. The members of this class attack the blood in various ways, some combine with the hæmoglobin and render it functionless as an oxygen carrier without otherwise interfering with the integrity of the red corpuscles: carbon monoxide is a typical poison of this group, to which hydrocyanic acid may be said to belong, together with sulphuretted hydrogen, although, in cases of poisoning by this gas in the human subject, the combination of sulphur with hæmoglobin has not yet been detected. A second group includes poisons which dissolve the stroma of the red corpuscles and set free the hæmoglobin: arsenuretted hydrogen is the best example of this group, which includes phallin—a toxalbumin derived from the fungus *Amanita phalloides*—and helvellic acid (*Helvella esculenta*), various glucosides as sapotoxin, and other saponins and also solamin may be included in this group, as they dissolve the red blood corpuscles when brought into direct contact with them: but, being difficult of absorption in the intestinal canal, they do not act as blood poisons when swallowed. A third group comprises a number of substances which convert the hæmoglobin: either whilst it is still within the corpuscles, or after setting it free into methæmoglobin, many members of this group are of no practical interest to the toxicologist. Among the more important are potassium chlorate, toluylendiamine—a typical example of those poisons which convert the hæmoglobin within the intact corpuscles into methæmoglobin, phenylhydroxylamine—a type of the poisons which disintegrate the red corpuscles and convert the liberated hæmoglobin into methæmoglobin, nitrobenzine, dimtrobenzine, anilin, with the countless coal-tar derivatives, which numerically increase year by year, as antifebrin, antipyrin, phenacetin, exalgin, pyrogallol, nitro-glycerine, amyl and other nitrites, picric acid, sulphonal, and trional. A fourth group is composed of substances which act differently on the blood to any of those already mentioned, they promote abnormal coagulation of the blood whilst it is within the vessels. The effect of some is to closely imitate physiological blood-coagulation: such

as ricin—a phytalbumose derived from the *Ricinus communis*, and abrin—a toxalbumin obtained from the *Abrus precatorius*, others impart to the red corpuscles a tendency to agglutinate together so that they form thrombi, as is the case in acute phosphorus poisoning. If added to freshly drawn blood, paraphenylenediamine produces immediate coagulation and converts the hæmoglobin either into methæmoglobin or into hæmatin, but when absorbed into the circulation its tendency to alter the blood is less powerful, although if directly injected into a vein coagulation at once takes place.

### EVIDENCE OF POISONING FROM THE DEAD BODY.

As the symptoms of poisoning during life vary according to the nature of the poison that has been taken, so do the post-mortem appearances differ when the result has been fatal. The two classes of poisons which yield the *most characteristic* post-mortem appearances are **corrosives** and **irritants**. All corrosives and all irritants do not produce the same after-death appearances, but certain characteristic indications are usually met with in each class respectively.

When making a post-mortem examination in a case of suspected poisoning, it is important to distinguish between the effects of **poison** and those due to **disease** on the one hand and to incipient **putrefaction** on the other. The post-mortem indications afforded by corrosive and irritant poisons are to be sought for along the digestive tract. They comprise *hyperæmia*, *softening*, and *ulceration* of the mucous membrane with *perforation* of the wall of a viscus either due to ulceration or, more frequently, to the direct action of a corrosive.

The ordinary effect of an irritant poison is to cause **hyperæmia** of the mucous membrane of the œsophagus, stomach, and possibly of the small intestine. The hyperæmia may either be diffuse, or isolated in patches, it is usually most marked about the cardiac end of the stomach, more rarely is the pyloric end affected. With some irritants there is a tendency to the formation of small hæmorrhagic points, or stræ, or there may be large dark patches, which stand out in contrast with the neighbouring and less deeply coloured mucous membrane. The hyperæmia may be most intense along the summits of the rugæ of the gastric or intestinal mucous membrane, or the entire mucous coat of the stomach may be hyperæmic and thickened, presenting a velvety appearance—frequently seen in acute arsenical poisoning. The mucous surface may be covered with a viscid secretion, which may be blood-stained.

**Softening** of the mucous membrane of the stomach when due to poison is usually caused by corrosives, indications of the same condition are to be found in the œsophagus and possibly in the mouth as well. It is not a common result of poisoning, when it occurs it is chiefly due to the direct chemical action of the poison. All corrosives do not cause softening, carbolic acid corrugates and hardens the mucous surfaces with which it comes in contact, and some other corrosives occasionally do the same. Softening is an almost invariable result of poisoning by the alkalis.

**Ulceration** of the mucous membrane of the stomach is occasionally seen as a result of irritant poisoning, it appears to be chiefly due to the local action of a portion of the poison on a limited surface of the membrane, in this way phosphorus may set up ulceration as a primary result, apart from its secondary effects. Sometimes ulceration is due to infarcts, the result of inflammatory processes causing blood-stasis. Removal of patches of mucous membrane by the direct action of corrosives is to be distinguished from ulceration—the

former is due to chemical destruction of tissue, the latter to pathological processes

**Perforation** as a result of poisoning is usually due to the direct chemical action of a corrosive on the coats of the stomach, or more rarely of the small intestine. The appearance of a perforation thus caused is characteristic: there is no indication of limitation by inflammatory processes, the margins, instead of being thickened, are partially disintegrated, a condition that extends some distance from the aperture, which is usually large and is irregular in outline, the edges of the opening and the contiguous parts (when the perforation is due to sulphuric acid) are blackened and charred. Ulceration due to irritant poisoning may be followed by perforation, but such an event is of exceptional occurrence, the appearances would more nearly resemble those met with when an idiopathic gastric ulcer has given way. The poison which most frequently causes perforation is strong sulphuric acid.

#### THE LOCAL EFFECTS OF POISON CONTRASTED WITH THOSE OF DISEASE AND WITH THOSE PRODUCED BY POST-MORTEM CHANGES.

Some of the above described appearances cannot be distinguished from similar appearances which are due to disease, others are only met with as the result of poisoning. Acute idiopathic inflammation of the mucous membrane of the stomach is exceedingly rare, so that the inflammatory appearances described when distinctly present, are always suspicious of irritant poisoning, on the other hand, such appearances may be absent, or feebly marked, after death from an irritant poison. The colour of the gastric mucous membrane should always be noted at the time the stomach is opened, as it becomes redder on exposure to air. The colour alone is *not to be relied on* as an indication of inflammation, it may result from food or medicine which contains pigmentary matter, some fruits, such as black cherries and elder-berries, will colour the gastric mucous membrane, which is also reddened during digestion of food and, it is said, by copious draughts of ice-cold water. Post-mortem staining produces redness, but it is limited to the posterior part of the stomach—the body having lain on its back, the mucous membrane is not thickened, there is no glairy mucous on its surface, and the general appearance is unlike that due to inflammation. Along with colour changes incipient **putrefaction** produces **softening** of the mucous membrane of the stomach commencing on the posterior part, at the seat of post-mortem staining, the softening affecting the entire thickness of the coats of the stomach. When softening is the result of **poisoning** it is either limited to the mucous membrane, or, if it extends to the muscular coat, the mucous membrane will probably be detached in patches over the parts that are softened, whilst in post-mortem softening the mucous membrane is rarely detached—the several coats of which the stomach is composed soften together usually without separating from each other.

**Ulcer** of the stomach is much more frequently due to pathological processes than to poisoning. The **idiopathic** gastric ulcer is small, sharply defined, and very frequently is situated along or near the lesser curvature. The floor of the ulcer is formed by the muscular coat or, if this is perforated, by the peritoneum, which, at the spot, may or may not be adherent to the liver or pancreas, the opening in the mucous membrane is circular and cleanly “punched out,” and

is larger than that through the muscular coat. In the early stage of the formation of a gastric ulcer the edges are not raised, subsequently they may become so, the mucous, muscular, and serous coats are firmly adherent for some distance from the ulcer. If the ulcer gives way a small opening usually forms through the floor, so that, in vertical section, the ulcer is V-shaped—the aperture being at the apex, all this points to gradual formation. An **ulcer** caused by an **irritant poison** is usually more quickly produced, and presents the appearance of an erosion with surrounding signs of recent inflammation, which are generally absent in the idiopathic ulcer, there is less tendency to thickening round the margin, which is more irregular and not so cleanly “punched out.” When perforation is due to the immediate action of a corrosive, the comparatively large size of the aperture, the irregular and ill-defined margin, the surrounding softening and friability of all the coats, and the discoloration of the structures make the diagnosis easy.

**Perforation** of the stomach wall may take place from the action of the gastric juice **after death**. This could not be mistaken for the effect of a corrosive poison, as there is entire absence of any indications of inflammation, and the edges of the aperture, though irregular, are free from the colour changes which characterise the action of corrosives, the surrounding mucous membrane is often swollen and gelatinous.

The appearance presented by the mucous membrane of the stomach in cases of suspected poisoning is to be *interpreted with caution*, especially in relation to colour-changes. Mere redness of the surface is too often assumed to indicate the occurrence of inflammation, which has at once attributed to the effects of poisoning, as previously stated, something beyond this is required to warrant such an interpretation.

The post-mortem appearances after death from narcotics, convulsives, and delirians, are mostly limited to hyperæmic conditions of the nervous centres and their membranes, and are relatively of little diagnostic value.

## THE SALE OF POISONS.

The Pharmacy Act of 1868 contains a **Schedule of Poisons**, the sale of which is restricted by various regulations. The schedule, which is divided into two parts, may be added to from time to time by the Council of the Pharmaceutical Society, with the approval of the Privy Council, and now stands as follows —

### PART I

**Aconite, Aconitine**, and their preparations

**Arsenic**, and its medicinal preparations (*see Note below*)

**Alkaloids**—all poisonous vegetable alkaloids not specifically named in this Schedule, and their salts, and all poisonous derivatives of vegetable alkaloids

**Atropine**, and its salts and their preparations.

**Belladonna**, and all preparations or admixtures (except belladonna plasters) containing 0.1 or more per cent of belladonna alkaloids

**Cantharides**, and its poisonous derivatives

**Coca**, any preparation or admixture of, containing 0.1 or more per cent of coca alkaloids

**Corrosive Sublimate**.

**Cyanide of Potassium**, and all poisonous cyanides and their preparations

**Diamorphine** (also known as Heroin) and all preparations or admixtures containing 0.1 per cent of Diamorphine

**Diethyl-Barbituric Acid**, and other alkyl, aryl, or metallic derivatives of barbituric acid, whether described as veronal, propanal, mednal, or by any other trade name, mark, or designation, and all poisonous urethanes and ureides.



- Eggonine**, and all preparations or admixtures containing 0.1 per cent of Eggonine
- Emetic Tartar**, and all preparations or admixtures containing 1 or more per cent of emetic tartar
- Ergot of Rye**, and preparations of ergots
- Lead** in combination with oleic acid, or other higher fatty acids, whether sold as diachylon or under any other designation (except machine spread plasters)
- Nux Vomica**, and all preparations or admixtures containing 0.2 or more per cent of strychnine
- Opium**, and all preparations or admixtures containing 0.2 or more per cent of morphine
- Picrotoxin**.
- Prussic Acid**, and all preparations or admixtures containing 0.1 or more per cent of prussic acid
- Savin**, and its oil, and all preparations or admixtures containing savin or its oil

*Note*—It is unlawful to sell arsenic (including arsenious acid, arsenites, arsenic acid, arseniates, and all other colourless preparations of arsenic), unless, in addition to the requirements of the Pharmacy Act, 1868, the provisions of the Arsenic Act be observed

## PART II

- Almonds, Essential Oil of** (unless deprived of prussic acid)
- Antimonial Wine**.
- Cantharides**, tincture and all vesicating liquid preparations or admixtures of
- Carbolic Acid**, and liquid preparations of carbolic acid and its homologues, containing more than 3 per cent of those substances, except preparations for use as sheep wash or for any other purpose in connection with agriculture or horticulture, contained in a closed vessel distinctly labelled with the word "Poisonous," the name and address of the seller, and a notice of the special purposes for which the preparations are intended
- Chloral Hydrate**
- Chloroform**, and all preparations or admixtures containing more than 20 per cent of chloroform
- Digitalis**
- Mercuric Iodide**.
- Mercuric Sulphocyanide**
- Oxalic Acid**.
- Poppies**, all preparations of, excepting red poppy petals and syrup of red poppies (*Papaver rhæas*)
- Precipitate, Red**, and all oxides of mercury
- Precipitate, White**.
- Strophanthus**.
- Sulphonal**, and its homologues, whether described as trional, tetronal, or by any other trade name, mark, or designation
- Zinc Chloride**, and liquid preparations of Zinc Chloride, except preparations intended to be used for soldering or other purely industrial purpose, provided that they are contained in closed vessels labelled with the word "Poisonous," and bearing the name and address of the seller and a notice of the special purpose for which the preparations are intended
- All Preparations or Admixtures** which are not included in Part I of this Schedule, and contain a poison within the meaning of the Pharmacy Act, except preparations or admixtures the exclusion of which from this Schedule is indicated by the words therein relating to carbolic acid, chloroform, and coca, and except such substances as come within the provisions of Section 5 of the Poisons and Pharmacy Act, 1908

It is important to notice that the effect of the last paragraph of Part II is to include in Part II many preparations and admixtures which are not specifically named in the schedule, including preparations and admixtures of non-scheduled vegetable drugs—such as Calabar bean, colchicum, conium, gelsenium, hyoscyamus, lobelia, stavesacre, stramonium, etc.—which contain poisonous alkaloids

Section 17 of the Act makes it unlawful to sell **any poison in either Part I. or Part II.** unless the box, bottle, vessel, wrapper, or cover in which the poison is contained is distinctly labelled with

- (1) The name of the article
- (2) The word "Poison," and
- (3) The name and address of the seller

The further regulations, **applying only to Part I.** of the schedule, make it unlawful to sell any poison in Part I to any person unknown to the seller unless introduced by some person known to the seller, and require that on every sale of such poison, the seller must before delivery enter in a book kept for the purpose, and termed the Poison Book

- (1) The date of sale
- (2) The name and address of the purchaser
- (3) The name and quantity of the article sold
- (4) The purpose for which it is required

These entries must be attested by the signatures of the purchaser and of his introducer if any

**Application to Medical Practitioners.**—By Section 3 of the Amending Act of 1869, the foregoing provisions are made not to apply to any medicine supplied by a legally qualified medical practitioner to his patient, provided such medicine be distinctly labelled with the name and address of the seller, and the ingredients thereof be entered with the name of the person to whom it is sold or delivered, in a book to be kept by the seller for that purpose

**The Arsenic Act, 1852,** requires that arsenic and its colourless preparations shall when sold be mixed with at least one-sixteenth its weight of soot, or half as much indigo, unless sold in a quantity of not less than ten pounds, and for a purpose (not for use in agriculture) for which such admixture would render it unfit

**Regulations for keeping, dispensing, and selling Poisons**—The following regulations have been prescribed by the Pharmaceutical Society with the consent of the Privy Council

1 That in the keeping of poisons each bottle, vessel, box, or package containing a poison be labelled with the name of the article, and also with some distinctive mark indicating that it contains poison

2 Also that in the keeping of poisons, each poison be kept on one or other of the following systems, viz —

(a) In a bottle or vessel tied over, capped, locked, or otherwise secured in a manner different from that in which bottles or vessels containing ordinary articles are secured in the same warehouse, shop, or dispensary, or

(b) In a bottle or vessel rendered distinguishable by touch from the bottles or vessels in which ordinary articles are kept in the same warehouse, shop, or dispensary, or

(c) In a bottle, vessel, box, or package kept in a room or cupboard set apart for dangerous articles

3 That in the dispensing and selling of poisons all liniments, embrocations, lotions, and liquid disinfectants containing poison be sent out in bottles rendered distinguishable by touch from ordinary medicine bottles, and that there also be affixed to each such bottle (in addition to the name of the article, and to any particular instructions for its use) a label giving notice that the contents of the bottle are not to be taken internally

**Sale of Poisonous Substances**—It is unlawful to sell any of the following substances unless the box, bottle, vessel, wrapper, or cover in which the substance is contained is distinctly labelled with the name of the substance, the words "Poisonous—Not to be Taken," and the name and address of the seller —Sulphuric, Nitric, and Hydrochloric Acids, Soluble Salts of Oxalic Acid, Liquid Preparations containing more than 5 per cent by weight of free Ammonia, and all Liquid Preparations sold as Carbolic, or Carbohc Acid, or Carbohc Substitutes, or Carbohc Disinfectants, containing not more than 3 per cent of phenols. Further, any liquid substance required to be so labelled may not be sold by retail except in bottles or other containers rendered distinguishable by touch from ordinary bottles or containers

**Dangerous Drugs Act.**—Under this Act regulations are made by the Home Office for controlling, and restricting the possession, sale, and distribution of raw opium, morphine, cocaine, ecgonine, and diamorphine (heroin)

### EVIDENCE OF POISONING FROM CHEMICAL ANALYSIS OF THE VISCERA AND THEIR CONTENTS.

The substances obtained at the post-mortem examination of a case of suspected poisoning are sent to the analyst in bottles, or jars duly secured and sealed, as directed when the method of making such examinations in medico-legal cases was described. Before opening the jars an inventory of them should be taken, and the covers and seals should be carefully scrutinised, in order to ascertain whether they have been tampered with, once in the possession of the analyst the jars and their contents must be kept under lock and key.

After each jar is opened its contents are to be measured, or weighed, in accordance as to whether they are fluid or solid. A careful examination of the physical appearance of the various substances should then be made, a lens or microscope being used if necessary. All food-stuffs present should be noted, and the odour of the contents of each jar ascertained, should any crystals or particles of inorganic matter be present, a few should be picked out and submitted to a preliminary examination, any seeds or fragments of leaves of plants that may be observed should be removed and investigated as to their nature and source.

When the symptoms observed during life are either obscure or are not indicative of a special poison, or in cases where no history is forthcoming, it may be necessary in the absence of post-mortem indications to make a systematic analysis of the viscera and their contents. Usually some clue is obtainable as to the nature of the poison suspected to have been administered, and in such cases the chemical investigation is chiefly directed to the discovery of that poison. The amount of material at the disposal of the analyst being limited, it is very important that he should be made acquainted, as far as possible, with the nature of the poison, as indicated by the symptoms during life and the appearances after death. It does not follow that the chemical enquiry is to be restricted to the discovery of a suspected poison: the possible presence of other poisons is not to be ignored. Usually, however, only one poison is present, and, if the analysis is from the first directed towards its discovery, the probability of success is much greater than if the investigation has to include the whole series of poisons. This is of special importance as regards quantitative analysis, and in criminal trials much importance is not unfrequently attached to the amount of the poison obtained from the dead body. Only a portion of the organs and substances at the disposal of the analyst should be used in making the analysis: unless the amount is too small, half should be reserved for a corroborative investigation by another analyst.

In making a **systematic analysis** attention is **first** to be directed to the possible presence of **volatile poisons**. The chief volatile poisons are—hydrocyanic acid, oil of bitter almonds, nicotine, conine, phosphorus, alcohol, chloroform, benzene, and its derivatives nitro-benzene, aniline, and phenol. If the odour of any of these bodies is present in the substance under investigation, the clue thus afforded is to be followed up, in any case their presence or absence must be determined.

The next step is to ascertain whether any **alkaloids** are present, for this purpose, one of the many modifications of Stas' process for the separation of alkaloids from organic matter may be resorted to. The principles on which this process is founded are—that the salts of the alkaloids are soluble in water and in ethyl alcohol, but not in ether and some other solvents, as amyl alcohol,

benzene, acetic ether, and chloroform, whilst, on the other hand, the uncombined alkaloids (or most of them) are nearly insoluble in water, but are more or less soluble in ether and in the other solvents named. This property of the alkaloids is made use of to extract them from organic admixture in the following way - The alkaloid present is dissolved with the aid of slightly acidulated alcohol, with which the viscera are digested for several hours at a moderate temperature, the liquid is then filtered off and at a gentle heat is evaporated down to a syrup. When cold, the syrupy mass is treated with absolute alcohol, the object being to precipitate as much of the foreign matter as possible, and to retain the alkaloid in solution, the process of evaporation and subsequent treatment with absolute alcohol may have to be repeated several times before the bulk of the extraneous matter is got rid of. Finally, the last alcoholic extract is evaporated down to a syrup, which is then dissolved in a small quantity of water, with the result that any alkaloid in the original substance will be held as a salt in aqueous solution. As long as this solution remains acid it may (except in the case of certain alkaloids and active principles) be shaken with ether without parting with the alkaloid, and by repeated shaking-out with ether, more of the remaining organic impurities, chiefly fatty matter, may be removed. When this is accomplished, the aqueous solution is made alkaline, and is once more shaken out with ether. The addition of the alkali displaces the acid of the alkaloid, and, being insoluble in water and soluble in ether, the free alkaloid is taken up by the ether, which is then separated and evaporated to dryness, leaving the alkaloid in a sufficiently pure state for testing.

So much for the principles on which Stas' process is founded, their successful application demands skill, and also attention to several important details. Stevenson's<sup>1</sup> great experience has enabled him to develop and to refine Stas' process as follows -

The substance under examination is digested with twice its weight, or, if fluid, twice its volume, of rectified spirit, at a temperature of 35° C, after several hours the fluid is poured off (the solid matter being subjected to pressure), and is replaced with fresh spirit, which is allowed to digest as before, after decantation of the second extract the process is repeated several times with spirit acidulated with acetic acid. The extracts obtained with the acidulated spirit are mixed together, but are kept apart from those obtained without the acid, which are also mixed together. The extracts separately are quickly raised to a temperature of 70° C, allowed to cool, filtered, and the residue on the filter is washed with spirit. The extracts are then evaporated to a syrup at a temperature not exceeding 35° C, excess of acid being neutralised with soda. The syrupy liquid is drrenched with 33 c.c. of absolute alcohol, and well stirred in a mortar, the alcohol is poured off, and the process is repeated with successive quantities of 15 c.c. of alcohol until it comes away colourless, these extracts are filtered and evaporated to a syrup as before. The syrupy extracts from the acid and from the non acid digestions are each diluted with a small quantity of water, filtered, and then mixed together. The united extracts, whilst still acid, are shaken with twice their volume of ether, the operation being repeated until, on evaporation of a few drops, the ether leaves no residue. The ethereal solutions are washed by vigorous shaking with 5 c.c. of water, to which a few drops of H<sub>2</sub>SO<sub>4</sub> have been added. The acid aqueous solution which was washed with the ether, and the water which was used to wash the ether after separation, are mixed, made alkaline with sodium carbonate, and then exhausted, first with a mixture of one volume of chloroform and three volumes of ether (which previously has been well washed with water), and subsequently twice or thrice more with washed ether alone. The ethereal extracts are washed with 5 c.c. of water, then with 10 c.c. of water acidulated with H<sub>2</sub>SO<sub>4</sub>, and again with 5 c.c. of water alone. The acid liquid and the final wash water are washed once or twice with a little ether, realkalified with sodium carbonate, and well extracted with washed chloroform and ether, and afterwards with ether alone. These ethereal extracts are washed with water barely alkaline with

<sup>1</sup> Watts' Dictionary of Chemistry, 1890

sodium carbonate, filtered through a dry filter, and evaporated under  $35^{\circ}\text{C}$  in tared glass basins in an oven. When evaporation is complete the basins may be dried at a temperature of  $100^{\circ}\text{C}$ , then cooled over sulphuric acid and weighed. To extract morphine Stevenson uses a well-washed mixture of equal volumes of acetic acid and ethylic ether.

The object of these repeated washings and transferring of the alkaloids from water to ether, and from ether to water, is to get rid of fatty and other matters which seriously interfere with the colour tests, and also to enable a correct estimation to be made of the amount of alkaloid present in the substance under examination. If the above-named temperatures are exceeded, some of the extraneous organic substances present become soluble in both water and alcohol, and are, therefore, extremely difficult to get rid of. When an alkaloid is sufficiently freed from organic matter the usual tests are applied to establish its identity.

When extracting an aqueous fluid containing an alkaloid with a fluid that is insoluble in water, care must be taken not to agitate the two fluids to such an extent as to cause them to emulsify, some fluids which retain small quantities of organic matter in solution very readily form emulsions with the solvent used for extraction, especially when alkaline. Various methods have been devised to promote separation of the fluids after emulsification, such as the addition of more of the solvent, the immersion of the containing tube for a few minutes in a freezing mixture, or in hot water, imparting a rotary motion to the tube, or a series of slight shocks by repeatedly tapping it with the finger nail—any of which may or may not be successful. The best plan is to avoid emulsification by first cautiously inverting the tube two or three consecutive times, and observing the rate at which the fluids separate, if they show a disposition to blend, the process of extraction must be conducted with great deliberation, time being allowed for separation after every two or three inversions of the tube.

The process of extraction may be conducted in a stoppered tube, like a large test tube, or in a small tubular separator, furnished with a stop cock, through which the lower stratum of fluid may be withdrawn. The separator is most convenient when the solvent is heavier than water, when a tube is used the solvent has to be pipetted off. The amount of solvent in toxicological work is usually small, and may be conveniently dealt with by means of a pipette furnished with an india rubber ball attached to its upper end by a short piece of rubber tube, on which is placed a spring pinch cock, as shown in Fig 21. Before using the pipette the pinch cock is opened, and the ball compressed so as to empty it of air, the pinch-cock is then allowed to close. The pipette is passed down to the lowest stratum of the fluid to be removed, and the pinch-cock gently pressed open, when the ball expands and draws up the fluid into the pipette. When the whole of the fluid, or as much as the pipette will contain, is removed, the pinch cock is again allowed to clip the tube, the pipette is withdrawn, and its contents are expelled by compressing the ball and opening the pinch-cock. The advantage of this device is that separation can readily be effected at the level of the eye, and arrested at the desired moment with the greatest nicety. The lower end of the pipette is turned sidewise, so as not to draw up the fluid beneath.

As regards modern synthetic drugs, Panzer<sup>1</sup> states that, with acid reaction, ether removes from aqueous solution, sulphonal, trional, veronal, hedonal, aspirin, salipyrin,

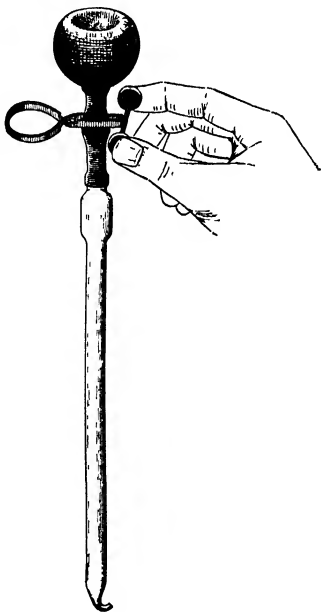


Fig 21

<sup>1</sup> *Ueber den forensischen Nachweis neuerer Arzneimittel*, 1906

and acetopyrin With alkaline reaction, ether removes pyramidon and antifebrin Amyl alcohol removes antipyrin and phenacetin

**Inorganic poisons** remain to be dealt with Mineral poisons maintain their integrity when submitted to processes that destroy alkaloids, therefore they can be treated in a different manner If inorganic poisons were simply mixed with organic matter it would be sufficient directly to convert them into insoluble salts, and to wash away the impurities the association between the two, however, is usually much more intimate, and then they can only be separated by destruction of the organic matter Some have to be separated from organic matter in special ways, or with special precautions, as they are liable to be dissipated by methods which do not affect most inorganic poisons, of these, arsenic is a notable example

Various processes are used to destroy the organic matter, of which three only need here be described, they are known amongst chemists as the moist and the dry methods The **moist method**, devised by Fresenius and von Babo, is accomplished as follows -

The substance which is suspected to contain the poison, if solid, is reduced to a pulp, and is mixed with sufficient water as to be of the consistence of thin gruel, bones should be divided into small fragments, urine should be evaporated to one fourth or one sixth its volume The substance thus prepared is placed in a large flask along with some crystals of potassium chlorate, each pound weight of the organic mixture will require about half an ounce of the chlorate Pure hydrochloric acid to about the same weight as the original substance is then added, and the flask is placed on a water bath and heated Chlorine, or rather a mixture of chlorine and of chlorine peroxide, is evolved, which attacks the organic matter, breaks it up, and liberates any mineral poison that may be present If necessary, additional crystals of potassium chlorate are added until the fluid becomes limpid and of a light yellow colour, or, if there is much organic matter, until it assumes the appearance and colour of thin oatmeal gruel On account of more gradual evolution of chlorine, the chlorate that is present before the flask is heated acts much more energetically, weight for weight, than fragments that are added after the liquid is hot, as a great deal of the gas then escapes without rendering any service Only moderate heat should be applied in order to minimise the waste of chlorine, and also the risk of frothing over—an accident very liable to occur, especially with substances containing sugar, starch, or alcohol After the last addition of chlorate, the liquid is transferred to an evaporating basin, and is allowed to remain on a water bath until the smell of chlorine has disappeared, it is then filtered hot The whole of the organic matter is not destroyed by this progress, fatty substances especially being resistant, but if the original matter is reduced to small fragments, any mineral poison present will be liberated

The objections raised against this process are that some important poisons, as arsenic and antimony, especially the former, are liable in part to escape in the form of vapour as chlorides, and that others, such as lead and silver, may remain as insoluble chlorides on the filter As regards the first objection, it is to be observed that when hydrochloric acid is diluted with water (as in the moist method of destroying organic matter) any arsenic which may be present in the hot solution is not given off with its acid aqueous vapour—arsenious chloride dissolved in hydrochloric acid being volatile only when the solvent is concentrated Any possibility of loss may be avoided, however, by furnishing the flask in which the organic matter is being destroyed with a condenser and receiver The second objection—as far as lead is concerned—is met by taking care to filter the solution whilst hot, if only a limited amount of lead is present it will remain in solution as chloride so long as the liquid is hot, and will consequently pass through the filter, a considerable quantity is kept in solution in the cold, as it forms a combination with potassium chloride, which is more soluble than lead chloride alone If a large amount is present it will not all be found in the filtrate, the substance left on the filter, therefore, should always be tested for lead, in toxicological work, however, the amount of lead present is not, as a rule, more than will remain dissolved in the cold Silver chloride, being insoluble either in hot or cold water, will not pass through the filter, consequently the salts of silver require dealing with in a special manner Arsenic also may be dealt with as described in the section devoted to that metal

Destruction of organic matter by the **dry method** is effected by heating it to redness so

that it is either carbonised or completely incinerated. When cold the residue is drenched with nitric acid, and sufficient heat is afterwards applied to drive off the free acid, the nitrate of the metal is then dissolved in water, filtered, and dealt with according to the kind of metal present.

The dry method is unsuitable in the case of the more volatile metals as arsenic, antimony, mercury, and, in a lesser degree, lead, tin, and zinc, further, it is exceedingly difficult and troublesome to carry out with large masses of organic matter, it is convenient with small amounts, and, in the absence of the more volatile metals, yields good results.

Another mode of destruction of organic matter is to heat it with an equal weight of  $\text{HNO}_3$  until the whole assumes the consistence of soup, then to add  $\text{KOH}$  till neutralised and to evaporate to dryness. The dry residue is deflagrated piece by piece in a porcelain capsule, additional saltpetre being added, if necessary, so as thoroughly to oxidise any metals present. The residue after deflagration is extracted with boiling water, the oxides of most of the metals that are insoluble in water will be rendered soluble by being treated with nitric acid. This method is convenient when operating on large masses of organic matter.

*The freedom from impurities of the chemical agents respectively used in these processes should invariably be ascertained.*

Special methods of separation will be described under the headings of the respective poisons

## INORGANIC POISONS.

## CHAPTER XXIX

## CORROSIVES.

## SULPHURIC ACID.

**Sulphuric Acid** ( $\text{H}_2\text{SO}_4$ ), or oil of vitriol, is a typical and most powerful corrosive. On coming in contact with organic matter, it combines with the water that may be present and chars the solids, if much water is present in the tissues, and the amount of acid is limited, they may be converted into a slimy mass of a brownish colour. Albumen is at once coagulated and subsequently dissolved. Muscle at first swells, becomes gelatinous, and then liquefies, the colour being brownish-red. Hæmoglobin is immediately converted into acid hæmatin. Sulphuric acid enters into chemical combination with albumen, and if no excess of acid is present, on digesting the albuminous compound with water, and applying the usual tests to the solution, no trace of free acid can be obtained. With progressive dilution of the acid the above-named effects diminish in degree until they cease to be produced.

**Symptoms.**—When the concentrated acid is swallowed intense pain is immediately experienced in the mouth, throat, and down to the stomach, from whence it rapidly spreads over the whole abdomen, the pain may be so violent as to cause tetanic spasms or general convulsions. Within a few minutes gaseous eructations, retching, and vomiting occur, the ejected matter consisting of coffee-coloured or blackish fluid (altered blood), with shreds and masses of mucous membrane and coagulated mucus. There is intense thirst with difficulty or impossibility of swallowing, each attempt being followed by renewed retching and vomiting. Respiration is laboured and noisy from tumefaction of the larynx. The voice is hoarse, or, probably, there is complete aphonia, and the mouth is filled with sticky mucus and shreds of membrane. The general condition is one of profound collapse—the skin is pale, cold, and clammy, that of the face being cyanosed from imperfect respiration, in some instances it is suffused and of a dusky red. The eyes are sunk and have a wild look, the pupils often being dilated. The pulse, of low tension, is quick and thready, frequently almost imperceptible. The bowels are almost invariably confined, in rare instances diarrhœa has occurred, the dejections containing altered blood and shreds of membrane. The urine is suppressed or nearly so, albumen, blood discs, hæmatin, and casts have been found in it, but their appearance is not constant, any sulphuric acid that has been absorbed is eliminated in combination, chiefly with calcium and as ether sulphates.

The mucous membrane of the mouth is swollen, corroded or excoriated,



the exposed parts being raw, sometimes the membrane is white, due to cloudy swelling of the epithelium. The lips are usually swollen and excoriated, and there may be further evidence of the action of the acid in the shape of brown streaks extending from the mucous lining to the skin on the lower jaw, especially at the angles of the mouth. In the case of young children the front part of the mouth may be free from corrosion on account of the acid having been administered in a spoon passed well back towards the throat.

Death may take place in the stage of collapse within a few hours after the poison is swallowed, if from the primary effects it is not likely to be delayed beyond twenty-four hours. Death is often sudden, either from asphyxia due to tumefaction of the glottis, or, probably, from pulmonary thrombosis or embolism caused by the action of the acid on the blood, or from perforation of the stomach. When death occurs *very* shortly after the poison is taken it may be due to shock.

If the patient survives the early stage, reaction sets in, the temperature rises and the pulse becomes fuller, the parts acted on by the acid slough and undergo the usual processes of separation, leaving a raw surface after they are detached. Several weeks after contact with the acid, a portion of the mucous membrane of the œsophagus may exfoliate and come away as a tube. Death from exhaustion may occur during the stage of reparation, towards the end of the first week being a fatal period. In some cases, during the second or third week, pain along the distribution of the intercostal and abdominal nerves has been observed, and in others diffuse hyperæsthesia, due probably to peripheral neuritis.

Should the strength of the patient hold out, cicatrization of the abraded parts begins and the raw surfaces gradually close in. This involves the usual result of the extensive formation of internal cicatrices—loss of a corresponding area of mucous membrane, and contraction of the newly formed tissue, which leads to stricture when a canal or an aperture is the part affected. The lower end of the œsophagus and the pylorus are the usual seats of stricture, less frequently the œsophagus is affected higher up, sometimes a stricture does not develop until the patient appears to have entirely recovered. The loss of gastric glands gives rise to *apepsia*, and the patient emaciates from insufficient nutrition due to this cause, or to the presence of an œsophageal stricture. Death may take place from inanition several months after the injuries were sustained.

**Fatal Dose.**—Half a teaspoonful of the strong acid killed a child a year old. One fluid drachm may be regarded as the smallest lethal dose for an adult, death at the end of a week having been thus caused in the case of a young man. Death has occurred within an hour, usually it takes place within thirty hours, but it may be indefinitely delayed when due to secondary causes. Recovery has taken place after an ounce of the strong acid had been swallowed.

Sulphuric acid has caused death by being accidentally administered as an enema, and also by being injected into the vagina in order to procure abortion.

The **prognosis** of sulphuric acid poisoning is unfavourable, from 60 to 70 per cent of the cases prove fatal.

**Treatment.**—Prompt neutralisation of the acid is the first step. For this purpose calcined magnesia is the best, the alkaline carbonates are not so good, but, since saving of time is all important, any alkaline substance at hand should be utilised. Egg-shells, chalk, whitening, or plaster chipped off a cornice or the ceiling, may be powdered and administered suspended in water. White of egg or soap and water are also handy remedies. If nothing else can be obtained

water should be freely administered. The stomach-tube must not be used. The next step is to mitigate pain by hypodermic injections of morphine. If the immediate effects are survived, nutrition will probably have to be maintained by nutrient enemata, preferably peptonised. Tracheotomy may be necessary.

**Post-mortem Appearances.**—The conditions found after death vary with the length of time the patient survived. If death takes place within twenty-four hours the lips will probably be corroded and stained brown, similar stains from spilling of the acid may be present on other parts of the surface. The clothing should be examined for indications of the action of the acid produced either in the act of administering or taking it, or by the early vomited matter, stained portions of fabric should be cut out and preserved for chemical examination. The buccal mucous membrane will be greyish, yellowish-white, or dark brown in colour, it will be softened and disorganised so as to be easily detached from the underlying structures, in parts it will probably be absent, the raw surface being covered with dark-coloured blood, a like condition extends to the pharynx and down the œsophagus, which appears contracted and thrown into longitudinal folds. The period of contract with the corrosive being shorter, the œsophagus is usually less profoundly affected than the stomach, exceptionally, the mouth and œsophagus may entirely escape injury. The author has seen, in the case of a child two years old who had swallowed a small quantity of sulphuric acid, a perforation of the fundus of the stomach as large as a shilling, with an extensive sharply defined surrounding area of corrosion and blackening, and yet neither the mouth nor the œsophagus showed the least trace of corrosion. The **stomach** is contracted, and on being opened shows evidence of profound disorganisation, any contents will probably consist of a viscid, dark-coloured substance (not necessarily having an acid reaction), which is blood partially changed into hæmatin mixed with serum and mucus. The gastric mucous membrane may be converted into a slimy coating, absent in parts, or it may be corrugated and hardened, in patches or strips it is dark coloured, even to being absolutely black, the areas which surround the parts chemically corroded by the acid show signs of intense inflammation. The entire coats of the stomach may be so far disorganised as to be easily torn, and there may be actual perforation, which occurs more frequently with sulphuric acid than with any other corrosive, the aperture is irregular, has blackened margins, and the coats of the stomach which form them are soft and friable. When the contents of the stomach escape through the perforation (which is not invariably the case) one or more of the neighbouring viscera may be corroded and even perforated from the outside—the colon has been thus perforated, and the surfaces of the liver and spleen have been rendered hard and friable. The corrosive action of the acid may extend beyond the stomach to the duodenum, in one case it could be traced to the ileum. It is noteworthy that the pylorus itself is usually only slightly attacked, the evidences of corrosive action frequently stopping suddenly short of the pylorus, even when a portion of the acid has passed through it into the duodenum. Fatty changes have been found in the liver and kidneys, the latter may yield indications of parenchymatous inflammation, hæmatin cylinders may occasionally be present in the tubules. The bladder is generally contracted and empty. Clots have been found in the blood-vessels, and the intra-vascular blood has been observed to be dark and tarry, or it may be converted into hæmatin, brown cylinders of which may extend from the stomach to the vena cava as far as the right side of the heart. Although the blood may be greatly reduced in alkalescence, it has never been found to yield an acid reaction during life, after death it may do so.

If the victim of sulphuric acid poisoning has survived a week or more, the post-mortem appearances will be modified accordingly. After a still longer interval, cicatrices will replace the corrosions, and the usual effects of stricture (if present) will be observed, if the lower end of the œsophagus is the seat of stricture, the part immediately above will be dilated, and the stomach probably contracted, in such cases the post-mortem appearances usually met with in cases of death from inanition will be more or less obvious.

**Chemical Analysis.**—In the examination of organic admixtures for sulphuric acid, it is first necessary to ascertain the presence of a free acid. This may be done by adding a few drops of the suspected fluid to a weak aqueous solution of tropæolin OO (diphenylamine-orange), which changes in the presence of free acid from light yellow to ruby or lake colour. This reagent reacts to a solution containing one drop of one of the mineral acids in 100 c.c. of water, and to a 0.5 per cent. solution of oxalic acid. It is not affected by the acid salts usually met with in toxicological investigations, it reacts, however, to such salts as potassium binoxalate and the bisulphates. Another way is to dissolve a fragment of the potassium-tartrate of iron of the Pharmacopœia in a little water so that it is tinged yellow, and to add a drop or two of a solution of potassium sulphocyanide, the addition of a liquid containing a free acid changes the colour to red. This reagent, less delicate than the preceding one, reacts to a solution containing four drops of a mineral acid in 100 c.c. of water, and to a 3 per cent. solution of oxalic acid. When the solution to be tested is feebly acid it should be trickled down the side of the test tube so as to float on the reagent, at the junction of the two fluids a red line is formed. Whilst resisting ordinary acid salts this reagent reacts to potassium binoxalate and the bisulphates.

If qualitative analysis only is aimed at, it will be sufficient if the suspected substance, after having been reduced by evaporation if necessary, is digested with alcohol and then filtered, the free acid is soluble, whilst the sulphates that may be present are insoluble in alcohol. The filtrate is neutralised with soda or potash, evaporated to dryness, and the residue, dissolved in water acidulated with hydrochloric acid, is subjected to the usual tests.

Although alcohol, by rejecting any combined sulphuric acid that may be present in the form of sulphates, prevents error as to the source of the acid which is obtained, it is open to the objection that a certain amount of the acid may enter into chemical combination with the alcohol, in which case the acid given up falls short of that which was originally dissolved.

When an exact estimation of the amount of free acid present in an organic mixture is required, advantage may be taken of the solubility of quinine sulphate in alcohol. Sufficient freshly precipitated quinine is added to the mixture as to take up the whole of the acid, the liquid is then evaporated to a paste and extracted with alcohol, which dissolves the quinine sulphate, but not any other sulphates that may be present. The alcoholic solution is filtered and evaporated to dryness, the residue being taken up by hot water, when cold, the quinine is precipitated as hydrate by the addition of ammonia water. The solution of ammonium sulphate thus formed is acidulated with hydrochloric acid and raised to 100° C., barium chloride is then added until the whole of the sulphuric acid is carried down as barium sulphate, the liquid being kept hot. The precipitate may be separated by filtration through a close filter (or, if preferred, by decantation), washed, and dried. Its weight multiplied by 411 equals the amount of concentrated sulphuric acid present. Barium sulphate will pass through almost any filter paper when precipitated in the cold, but by boiling it becomes granular, and can then be kept back by a close textured paper.

**Tests.**—A solution of barium chloride produces a precipitate of barium sulphate which is insoluble in hydrochloric acid. If some of the precipitate is mixed with an equal bulk of sodium carbonate and made into a paste with a few drops of water, and is then fused on charcoal with the aid of the blow-pipe, it is converted into a sulphide, and, when cold, may be submitted to the usual tests, a small fragment placed on a clean silver coin and moistened with water produces a brown stain of silver sulphide. If a drop of a liquid which contains free sulphuric acid is allowed to fall on a piece of filter paper which is then dried before the fire, the paper, where covered by the drop, becomes charred.

The presence of free sulphuric acid cannot always be demonstrated in the tissues of those who have succumbed to it. In the case of *Reg v Berry* (Liverpool Assizes, 1887), the prisoner was convicted of having poisoned her daughter, a young child—as *Harris*<sup>1</sup> and other medical men strongly suspected—with sulphuric acid. The usual indications of corrosive action in the lips, mouth, and œsophagus were present, and the stomach and small intestines were inflamed, but not corroded, no trace of the acid was found in the body. Cases have occurred in which the presence of sulphuric acid was readily demonstrated in the early vomit which fell on the clothing or floor, but after the death of the victim none could be found in the body. In one case, within an hour or two after half an ounce of oil of vitriol was swallowed, the mucous fluid that welled up into the mouth would not redden litmus paper, this disappearance of free acid is due to the combinations which it forms with basic substances contained in the organism.

### NITRIC ACID.

**Nitric Acid** ( $\text{HNO}_3$ ), or aqua fortis, is a powerful corrosive which produces symptoms resembling those caused by sulphuric acid. The chief points of difference are due to the fumes which are given off by the concentrated acid, to the colour of the stains produced by it, to the absence of charring of the tissues, and consequently to diminished liability to perforation of the stomach.

The **symptoms** come on immediately after the concentrated acid is swallowed. They comprise intense pain, gaseous eructations, retching, vomiting, and collapse. On account of the formation of a larger amount of gas, the abdomen is usually more distended, and, if possible, is more exquisitely tender than in sulphuric acid poisoning. The lips, tongue, and mucous membrane of the mouth are softened and swollen, and are of a yellow hue, resulting from the formation of xanthoproteic acid. The teeth are sometimes attacked, the acid may dissolve the enamel and colour them yellow. The air passages are more liable to be attacked than in the case with sulphuric acid, pneumonia being a very possible complication due to inhalation of the fumes of the acid. The remaining symptoms do not materially differ from those met with in sulphuric acid poisoning.

**Fatal Dose.**—The smallest recorded fatal dose is two drachms. Recovery has taken place after a dose of half an ounce. Death has occurred in less than two hours, but this is unusual, from twelve to twenty-four hours or more is the average duration of life in fatal cases.

**Treatment.**—As in sulphuric acid poisoning.

**Post-mortem Appearances.**—In acute cases, allowing for the difference in colour of the parts acted on and a somewhat less intensive corrosive action, the appearances resemble those produced by sulphuric acid. The mouth, teeth, and œsophagus present a tint varying from yellow to brown. The yellow stains due to nitric acid may be distinguished from those due to iodine by touching them with ammonia water—the colour remains or is deepened, if caused by iodine it disappears. The mucous membrane acted on is softened and easily detached, if the acid has reached the **stomach** its mucous coat is stained yellow in parts, there may also be patches of a blackish-brown, resulting from alteration of effused blood, and within the vessels masses of hæmatin may be present. The stomach may be perforated, but not so commonly as with sulphuric acid, if not perforated, the entire thickness of its wall may in parts be softened and friable. The duodenum may be affected in the same way,

<sup>1</sup> *Med. Chron.*, 1887

or both it and the stomach may be simply inflamed. When life has been prolonged for weeks or months, cicatrices and stricture may be present.

**Chemical Analysis—Tests.**—When mixed with organic matter (after proving the presence of a free acid), simple qualitative analysis may be made by neutralising the mixture with potassium carbonate, filter paper dipped in the solution and dried forms touch-paper, which deflagrates on ignition. On evaporating a little of the solution to dryness and adding a few drops of strong sulphuric acid to the residue, and then stirring in a crystal of brucine, a bright-red colour is produced, this test is very delicate, the reaction being sufficiently distinctive as to yield decisive results notwithstanding the presence of a considerable amount of foreign colouring-matter. If a crystal of ferrous sulphate is substituted for the brucine it becomes surrounded by a brownish ring, this test is valueless unless the solution to be tested is free from colour. If a fragment of gold leaf is boiled in a test-tube with a little strong hydrochloric acid, and a few drops of a solution containing nitric acid are added, the gold leaf is partially or wholly dissolved, the solution of gold chloride thus obtained should be proved to be such by the addition of a little stannous chloride, which produces the colour known as the purple of Cassius. If to an aqueous solution of diphenylamine a few drops of a liquid containing nitric acid or a nitrate are added, and subsequently a little concentrated sulphuric acid is gently poured down the side of the inclined test-tube, so as to form a layer at the bottom, a blue ring develops above it. Before using this test a control experiment should be made by adding sulphuric acid in the manner above described to the solution of diphenylamine, without any of the suspected fluid, as some specimens of sulphuric acid being contaminated with nitric or nitrous acids give the reaction alone. The test is so delicate that it reacts to a cubic centimetre of water which contains one drop of nitric acid to 100 c c.

For quantitative estimation when in organic admixture, the acid may be taken up by freshly precipitated quinine, the solution evaporated to a paste, and the paste extracted with alcohol. The alcoholic solution is then filtered and evaporated to dryness, the residue is dissolved in water and the quinine is precipitated with sodium hydroxide. The solution of sodium nitrate thus obtained is evaporated to a syrup, and is treated in a closed vessel for some time with powdered aluminium, or with nascent hydrogen evolved by a voltaic couple. It is then distilled into a receiver containing strong hydrochloric acid, excess of platinum chloride is added to the distillate, and the whole is evaporated to dryness. After being washed with small quantities of alcohol, and dried, the residue—ammonio platinum chloride—is weighed. 100 parts correspond to 28.3 parts of nitric acid. This method is founded on the power possessed by nascent hydrogen of converting the nitrogen of nitric acid into ammonia, which is estimated as the double chloride of ammonium and platinum.

**Nitric Acid fumes** have occasioned death in several instances. One of the masters and the janitor of the Edinburgh Institution were carrying a jar of nitric acid when it fell and broke, and, in endeavouring to save some of the spilt acid, they were exposed to the fumes. The master went home unconscious that anything was wrong with him, in an hour or two difficulty in breathing set in, and he died ten hours after the accident, the janitor died the following day.<sup>1</sup> Stickler<sup>2</sup> records an instance in which a bottle of nitric acid was broken in the hold of a vessel, several men went below to remedy the mischief, and felt no immediate trouble, but within a few hours they began to complain of illness, and in a short time died. Two firemen whilst helping to extinguish a fire in a chemical store inhaled the fumes of nitric acid, and died the same day.<sup>3</sup> A similar accident, recorded by Kunne,<sup>4</sup> happened to thirteen firemen

<sup>1</sup> *The Lancet*, 1863.

<sup>2</sup> *New York Med. Rec.*, 1886.

<sup>3</sup> *Pharm. Journ.*, 1890-91.

<sup>4</sup> *Deutsche med. Wochenschr.*, 1897.

who extinguished a fire in a building in which a large number of carboys filled with fuming nitric acid were stored, immediate tightness in the chest with an irritable cough, vomiting, and headache were experienced, the symptoms quickly passed off and the men went home feeling quite well. About six hours after, severe symptoms suddenly developed—difficulty of breathing, vomiting, rapid pulse (120), which subsequently became slow, cyanosis, insensibility, and cramps, with great restlessness. Two of the men died in a few hours, the remaining eleven recovered. Fine crepitations were heard at the front of the chest, and in some instances the expectoration was mixed with blood, in none of these cases was there any tendency to bronchitis.

It is noteworthy that in cases of poisoning by the fumes of nitric acid the sufferer feels nothing wrong for two or three hours after the primary irritation has passed off, dangerous symptoms then begin and, as a rule, death rapidly ensues. Occasionally recovery takes place after a prolonged period of extreme danger to life, the initial symptoms having been only moderately severe. Two men were admitted into hospital under the author's care, suffering from the inhalation of nitrous fumes liberated by the breaking of a large bottle of nitric acid in their workshop. On admission one of the men was extremely ill and cyanosed, the next day he was much better, and within a week was able to return home. The other man appeared to be only slightly affected on the first day, but became much worse on the following days, and hovered between life and death for upwards of a week, eventually he slowly recovered. In both these cases the blood was examined spectroscopically, but no abnormality was observed. Death is usually preceded by symptoms resembling those of pneumonia, or of capillary bronchitis, and the air-passages have been found blocked by softened membrane and mucus tinged with blood. In addition to the action of the acid vapour on the mucous lining of the bronchi and the air-vesicles, it appears that the pulmonary terminals of the vagi are rendered functionless, and the reflex of the respiratory centre is thus cut off, probably the pulmonary vaso-motor nerves from the sympathetic are also paralysed, the result is rapid excessive secretion of mucus into the bronchi and alveoli, with lessened capacity for its expulsion, and consequent speedy death from asphyxia. The formation of nitro-compounds with hæmoglobin has been assumed, but their existence is doubtful, conversion of part of the hæmoglobin into methæmoglobin has also been postulated. Schmeiden<sup>1</sup> examined the blood in a case of fatal poisoning by nitrous fumes, both before and after death, without discovering any abnormality in the spectrum.

When a large quantity of nitric acid is accidentally spilt, chalk should be thrown on it, and ventilation promoted, the individuals present keeping to the windward, and, as far as possible, holding their breath when near the acid.

### HYDROCHLORIC ACID.

**Hydrochloric Acid** (HCl), or spirits of salts, is a corrosive of less activity than the two preceding acids, but on account of its volatility it is more apt to attack the air-passages than either of them. The **symptoms** resemble, but are not so severe as, those produced by sulphuric acid. On account of its comparatively feeble corrosive action, hydrochloric acid scarcely stains the skin, if at all, thus greatly differing from the other mineral acids. In 1919, there were 11 accidental and 40 suicidal deaths in England and Wales from poisoning by hydrochloric acid.

<sup>1</sup> *Centralb f die ges. Med.*, 1892

**Fatal Dose.**—The smallest quantity that has proved fatal is one teaspoonful. In two instances, both young girls, this was sufficient to cause death, and in one (Von Beyerlein<sup>1</sup>) perforation of the stomach was produced. On the other hand, recovery has taken place after an ounce and a half of the commercial acid had been taken, calcined magnesia having been administered ten minutes after it was swallowed (Ross<sup>2</sup>). In another case, recovery took place after swallowing two ounces. Death has occurred in two hours, and has been delayed for several days, the usual period is from eighteen to thirty hours.

**Treatment.**—As for the other mineral acids.

**Post-mortem Appearances.**—The mucous surfaces acted on are usually of an ashy-grey colour, interspersed with erosions. The inner surface of the stomach may appear reddened from acute gastritis, or blackened at parts from extravasation of blood which has been acted on by the acid. Perforation is exceptional, but, as shown in the case above mentioned, may occur even with a minimum fatal dose, in other respects the post-mortem appearances resemble those seen in sulphuric acid poisoning, but they are less pronounced. When a large quantity of the strong acid is swallowed, extensive disorganisation of the viscera may result. A young girl swallowed one hundred grammes (three fluid ounces) of hydrochloric acid, and died nine hours afterwards, at the autopsy a large perforation of the stomach was found, and the peritoneum and the liver were disorganised by the extravasated acid, which had also perforated the diaphragm and attacked the left lung (Burdett<sup>3</sup>).

**Chemical Analysis**—After ascertaining the presence of a free acid, as described when dealing with sulphuric acid, an organic admixture containing hydrochloric acid may be distilled, and the distillate tested qualitatively and quantitatively.

**Tests.**—When silver nitrate is added it produces a precipitate of silver chloride, which is insoluble in nitric acid, but is soluble in ammonia water. A fragment of gold leaf in boiling nitric acid is dissolved if a few drops of the solution containing hydrochloric acid are added. The amount of hydrochloric acid present may be estimated by precipitation, as silver chloride, which is dried, ignited, and weighed, 100 parts correspond to 25.44 parts of hydrochloric acid. If preferred, the amount of acid present in the distillate may be estimated volumetrically.

## OXALIC ACID.

**Oxalic Acid** ( $\text{H}_2\text{C}_2\text{O}_4 \cdot 2\text{H}_2\text{O}$ ), in combination with soda, potash, and lime, occurs in many plants and vegetables, as sorrel, rhubarb, the common dock, and some lichens. The acid is used commercially in the manufacture of straw mats, for the cleansing of articles of brass, in dye- and print-works, and also for domestic cleansing purposes. It is soluble in about 10 parts of cold water, and in about  $2\frac{1}{2}$  parts of alcohol. It volatilises at a temperature of  $150^\circ$  without residue, and in this way may be distinguished from its combinations (except with ammonia), which leave residues. When heated with strong sulphuric acid, it is decomposed without blackening, into water, carbon dioxide, and carbon monoxide. Oxalic acid crystallises in the form of slender prisms, and on this account has been mistaken for Epsom salts.

When swallowed in poisonous doses, oxalic acid produces local effects which resemble those produced by the mineral acids, it also exercises a special influence on the nervous system and upon the action of the heart.

<sup>1</sup> *Friedreich's Blätter f. ger. Med.*, 1890.

<sup>2</sup> *The Lancet*, 1886.

<sup>3</sup> *Lyon Medical*, 1895.

**Symptoms.**—The symptoms vary not only with the amount of acid, but also with the concentration of the solution in which it is taken. If half an ounce or more of the acid is dissolved in water so as to form a concentrated solution, the local effects resemble those produced by the mineral acids. Immediately, or soon after the solution is swallowed, pain is felt in the mouth and throat, which extends to the stomach and radiates over the abdomen, vomiting usually quickly follows and persists, the vomited matter consisting chiefly of altered blood. Diarrhœa is exceptional. Benjamin<sup>1</sup> states that in twenty-seven consecutive cases it scarcely ever occurred. If the acid is swallowed in more dilute solution, the above-named symptoms are delayed and are less violent. General indications of collapse are manifest, the respiration is gasping, the pulse small and irregular, and the extremities, or even the entire surface, may be cyanosed. Clonic spasms are not unfrequent, they may alternate with tonic contractions of the muscles, especially of those of the lower jaw, producing trismus. Aphonia may occur, and may persist for some time during convalescence.

The effects produced by oxalic acid on the nervous system are exceedingly irregular, sometimes they constitute the chief symptom, and at others they do not exceed those which might be reasonably expected from the reflex of the part with which the poison comes directly in contact. The nerve symptoms comprise paræsthesiæ and anæsthesia of the limbs or trunk, with aching and shooting pains in the loins, numbness of the tips of the fingers has been observed, and tenderness of the muscles of the legs, convulsions are common, and sometimes resemble those due to strychnine. Oliver<sup>2</sup> relates a case in which a poisonous dose of oxalic acid produced muscular tremor with greatly exaggerated knee-jerk. While the patient lay quietly in bed, sudden contraction of the diaphragm repeatedly occurred, producing an abrupt, deep, prolonged inspiration, and occasionally a snorting expiration. In other cases oxalic acid seems to act as a narcotic. The author saw a case of this kind in which the sufferer lay unconscious, breathing stertorously, the surface being cold and clammy as in opium poisoning, without there being any vomiting or other signs of gastric irritation. When the nervous symptoms form the prominent feature of the case, the poison has often been taken well diluted, or the stomach has contained some amount of solid food, the indications of gastritis being slight or entirely absent, in these cases post-mortem signs of local irritation, although present, are not as well marked as is usual. Large crystals of calcium oxalate are frequently present in the urine.

In 1919, there were 2 accidental and 37 suicidal deaths from poisoning by oxalic acid in England and Wales.

**Fatal Dose.**—The smallest recorded fatal dose is 60 grains, which taken in the solid form caused the death of a boy aged sixteen. Recovery has occurred after an ounce and a quarter. Death has occurred in ten minutes, usually it takes place within two hours, but it may be delayed for several days—seven in one case, and twenty-one in another.

**Treatment.**—Chalk or whiting suspended in a small quantity of water or milk may be given, although the consequent liberation of carbonic acid is disadvantageous, saccharated solution of lime has been recommended by Husemann. Plaster chipped from the walls, or egg-shells powdered and suspended in a little water are good, calcined magnesia may be given. The alkalis or their carbonates should not be given, as the resulting compounds are both soluble and poisonous. All antidotes should be given in as little water

<sup>1</sup> *Charité-Annalen*, 1899.

<sup>2</sup> *Brit. Med. Journ.*, 1895.



as possible in order to limit diffusion of the poison, its action not being solely local. After the acid is neutralised the bowels should be relieved by an enema or by castor oil.

**Post-mortem Appearances.**—These vary in accordance with the amount of the poison and the concentration of the solution, if the poison is taken in concentrated solution, or in the solid form, the local effects will probably be well marked. There may be corrosion of the mucous membrane of the mouth, œsophagus and stomach, or it may be white, softened, and easily detached from its bed. The inner surface of the œsophagus may be longitudinally corrugated, displaying numerous small erosions. The degree of inflammation of both œsophagus and stomach varies from a slight redness to an almost gangrenous condition, the inflammation may reach the duodenum. In a specimen in the museum of Owens College, Manchester, the interior of the stomach is blackened, resembling the condition met with in poisoning by sulphuric acid. Perforation of the stomach is exceptional, although the walls are often considerably softened, the mucous membrane not unfrequently presents cloudy spots, which are due to deposition of calcium oxalate, they often occur near hæmorrhagic infarcts. The kidneys may show a whitish zone between the cortical and the medullary portions, due to the deposit of crystals of calcium oxalate chiefly in the convoluted, and to a lesser extent in the straight, tubules, the glomeruli are free from deposit (Kobert and Kussner<sup>1</sup>). Examined microscopically the deposit is seen to consist of rhombic prisms, or of octahedral crystals.

**Chemical Analysis—Tests.**—Oxalic acid forms a white precipitate of calcium oxalate on the addition of a solution of calcium chloride or calcium sulphate, the delicacy of the reaction being enhanced if the solution of oxalic acid is neutralised with ammonia before adding the reagent, calcium oxalate is insoluble in acetic acid, and is soluble in hydrochloric acid. Silver nitrate gives a white precipitate of silver oxalate, which is easily soluble in nitric acid, and also in ammonia. Lead acetate gives a white precipitate, which is soluble in nitric acid.

**Organic admixtures** may be evaporated down at a gentle heat and exhausted with hot alcohol, to which a little hydrochloric acid has been added after filtration, the alcoholic solution is evaporated to dryness, and the residue dissolved in water.

**Quantitative Estimation.**—To a measured quantity of the aqueous solution, calcium acetate is added in slight excess, and the deposit of calcium oxalate is separated, washed with acetic acid and then with water, and dried. By careful ignition at a moderate temperature the calcium oxalate is converted into carbonate. 100 parts correspond to 126 parts of crystallised oxalic acid.

**Potassium Binoxalate** ( $\text{KHC}_2\text{O}_4 \cdot 2\text{H}_2\text{O}$ ), or salt of sorrel, or, as it is also called, salts of lemon, is an acid salt soluble in 40 parts of cold and in 6 parts of boiling water. It is used in the household for removing iron stains from underclothing. It is nearly, if not equally, as poisonous as oxalic acid, like symptoms being produced. Half an ounce has proved fatal, Park<sup>2</sup> records a case in which although severe symptoms were produced, recovery took place after a like quantity. Braithwaite<sup>3</sup> records the case of a woman aged twenty-four who, after swallowing about three-quarters of an ounce of salts of lemon, died in twenty-five minutes. At the autopsy the lips and mouth were found to be corroded, and two perforations were found in the cardiac end of the stomach.

<sup>1</sup> Virchow's *Arch.*, Bd 78

<sup>2</sup> *The Glas. Med. Journ.*, 1889

<sup>3</sup> *Brit. Med. Journ.*, 1905.

The chemical tests are the same as for oxalic acid, from which it may be distinguished by the deposit of potassium carbonate which is left when potassium binoxalate is ignited on a slip of platinum foil

Two cases of supposed fatal poisoning by *Sorrel-leaves* in children are recorded by Flemyng<sup>1</sup>

### ACETIC ACID.

**Acetic Acid** ( $C_2H_3O_2$ ) in the concentrated form—known as glacial acetic acid—is a corrosive, when dilute it is simply an irritant. On account of its volatility and pungency, some of the acid, when swallowed, is usually aspirated into the larynx, hence severe asphyxial symptoms almost invariably occur.

**Symptoms**—The mucous membrane of the lips, mouth, tongue, and other parts with which the acid comes in contact, is softened and presents the appearance of having received a white or pale yellow coating. As with other corrosives, the front part of the mouth may occasionally escape. Luff<sup>2</sup> mentions the case of a young woman whose lips and tongue presented no appearance of corrosion although she had swallowed two teaspoonfuls of glacial acetic acid, in the act of swallowing the poison had probably been thrown to the back of the mouth. If the patient be seen early the odour of acetic acid will probably be perceptible in the breath. The general symptoms comprise pain from the mouth down to the stomach, vomiting, collapse, laboured noisy breathing, irritable cough, and, in some cases, tetanic spasms, of reflex origin, caused by the excessive pain. Bojasinski<sup>3</sup> saw four children who had been poisoned with acetic acid, tetanic spasms occurred, the symptoms being altogether of a suffocative type, resembling those of acute laryngitis. Tufanow<sup>4</sup> relates four cases of poisoning by acetic acid (80 per cent), three of which were fatal, the chief post mortem appearances were corrosion of the œsophagus, stomach, and intestines, with icterus and subendocardial hæmorrhages. Stumpf<sup>5</sup> saw a man aged thirty-two who took a tablespoonful of 50 per cent solution of acetic acid mixed with an equal quantity of water. Severe gastro intestinal irritation was produced, and death took place in three days. Post mortem, the mucous membrane of the stomach was dark grey in colour, and ecchymoses were present both in the stomach and duodenum. Gesslewitsch<sup>6</sup> reports the case of a man who took an unknown quantity of acetic acid with the intention of committing suicide. On the seventh day he vomited a cast 26 cm long consisting of mucous membrane, submucosa and part of the muscular coat. Gastrostomy was performed four weeks after the acid was taken, but death occurred three weeks later from exhaustion.

**The Fatal Dose**—One teaspoonful of glacial acetic acid caused the death of a child, and recovery has taken place in an adult after six fluid ounces of 80 per cent acid.

**Treatment**—The acid should be neutralised with magnesia and the resulting compound evacuated by procuring emesis if it does not occur spontaneously, subsequently opium may be advisable. The laryngeal symptoms may be alleviated by the application of ice compresses to the throat and by allowing the patient to suck small pieces of ice, or to swallow ice cold water. Tracheotomy may be necessary.

**Post-mortem Appearances**—In addition to the local appearances already named, a more or less similar condition will probably be found in the œsophagus and stomach. In the case of an infant ten months old, which was fatally poisoned with a teaspoonful of glacial acetic acid, Luff<sup>7</sup> found that the mucous membrane of the mouth, epiglottis, larynx, œsophagus, and stomach was eroded in patches, and that, in parts, the mucous membrane of the stomach was corrugated and congested.

**Chemical Analysis**—Free acetic acid may be separated from organic matter by distillation, if combined it should first be liberated by the addition of phosphoric acid.

**Tests**—Acetic acid may be recognised by its odour. With ferric chloride and ammonia to neutralisation, a red colour is produced which changes to yellow on the addition of hydrochloric acid.

### TARTARIC ACID.

**Tartaric Acid** ( $C_4H_6O_6$ ), on one or two occasions, has produced severe and even fatal poisoning. Chabrie<sup>8</sup> found that the toxic action of lævo-tartaric acid is twice that of the dextro acid, and that, with equal weights of acid, the strength of the solution in which it is administered considerably influences the result.

<sup>1</sup> *The Lancet*, 1896

<sup>2</sup> *Text-Book of Forensic Med. and Toxicol.*, 1895

<sup>3</sup> *Medycyna*, Warszawa, 1892

<sup>4</sup> *Congrès Internat. Méd.*, Moscow, 1897

<sup>5</sup> *Munchener med. Wochenschr.*, 1898

<sup>6</sup> *Petersb. Med. Ztschr.*, No. 1, 1914

<sup>7</sup> *Loc. cit.*

<sup>8</sup> *Comptes rendus de l'Acad. des Sc.*, 1893.

The **Symptoms**, as seen in the following case recorded by Trevithick,<sup>1</sup> are those of a strong irritant rather than of a true corrosive. A woman, aged 67, swallowed at least two teaspoonfuls of tartaric acid made into a strong solution, acute abdominal pain and vomiting quickly occurred, followed in a few hours by diarrhoea, death, from exhaustion, took place on the seventh day. At the autopsy there were indications of general peritonitis with subserous hæmorrhages on the stomach, erosions were found in the œsophagus, and the mucous membrane of the entire intestinal tract was inflamed.

The **Treatment** would be to neutralise the poison with alkalies and to give astringents and opium.

## POTASSIUM.

**Potassium Hydroxide (KOH)** is met with as a poison in the impure condition in which it is used in the arts and manufactures. It has a strongly caustic action on the tissues, and on account of its affinity for water the effects produced by it tend to radiate to a considerable distance from the spot to which it is applied. It combines with fatty matters and decomposes the soft structures, leaving a greasy mass which keeps soft and moist, the local damage thus differing from that produced by the mineral acids. In the form of a basic carbonate—as pearl ash—it acts much in the same way.

**Symptoms.**—When a strong solution of potash is swallowed an immediate burning sensation is experienced in the mouth and throat, which extends to the stomach and radiates over the abdomen. Vomiting usually occurs, the vomited matter has a strong alkaline reaction, is slimy and may be coloured by blood which escapes from the inner surface of the stomach and œsophagus, shreds of mucous membrane are generally present in the vomit. Purging is not unfrequent. Collapse quickly occurs, the pulse is thin and feeble, and the surface is cold and clammy. The lips, tongue, and inside of the mouth are red and swollen. Convulsions may occur as in sulphuric acid poisoning. Stricture of the œsophagus is very liable to follow in cases in which the immediate symptoms are recovered from.

**Fatal Dose.**—The smallest recorded fatal dose is 40 grains, usually a much larger dose would be required, probably 3 or 4 drachms. Death has occurred within a few hours, more frequently it results after weeks or months from secondary symptoms.

**Treatment.**—Vegetable acids, as acetic (vinegar) or citric (lemon juice), in a dilute form, should be given, with olive oil, demulcents, and opium. The stomach tube must not be used.

**Post-mortem Appearances.**—The lips and, possibly, the surrounding skin on the face may show traces of the caustic action of the poison. Within the mouth the mucous membrane will be softened and of a brown colour, in parts it will probably be detached. In recent cases the tongue will be swollen and inflamed, the mucous membrane of the pharynx and œsophagus presenting more or less the same appearance. The mucous coat of the **stomach** is inflamed and softened, the colour is not constant, sometimes it is bright red and at others dark, it may be completely or merely superficially eroded in parts. If, as is frequently the case, the patient survives some weeks, stricture is usually found at the lower end of the œsophagus or at the pylorus.

**Chemical Analysis.**—The organic substances containing potash may be evaporated to dryness and incinerated to burn off the organic matter. The residue is then dissolved in a small quantity of water, slightly acidulated with hydrochloric acid, and precipitated with platinum chloride, precipitation may be aided by the addition of alcohol. The precipitate, dried and then washed with alcohol in small quantities until it comes away colourless, is again dried and weighed, 100 parts correspond to 19.272 parts of potash.

<sup>1</sup> *Brit Med Journ*, 1893

**Tests.**—If there is much potash present in the original solution it may be directly precipitated as a double salt with platonic chloride. A saturated solution of tartaric acid also precipitates potash. Before the suspected substance is submitted to analysis, its alkaline reaction should be ascertained. The spectroscopic reaction of potash is too delicate to be of much use, as it demonstrates the presence of the alkali in the tissues and in articles of food that may accidentally be present.

### SODIUM.

**Sodium Hydroxide** (NaOH), or caustic soda, is largely used in manufactures, but cases of poisoning, which are usually accidental, are not frequent.

The symptoms, lethal dose, treatment, and post-mortem appearances are precisely the same as in poisoning with potash.

**Chemical Analysis**—For toxicological purposes there is no satisfactory chemical test for soda, it is best identified by exclusion. If there is a solid residuum after evaporation and incineration of the suspected substance, and both it and the original substance yield a marked alkaline reaction, and no precipitate is formed on the addition of platonic chloride to a concentrated solution of the incinerated product, the alkali which is proved to be present will be soda. Absence of the alkaline earths must be ascertained. The spectroscopic reaction of sodium is useless in toxicological investigations on account of the ubiquitous presence of its salts.

### AMMONIA.

**Ammonia Water** ( $\text{NH}_4\text{OH}$ ), also known as spirits of hartshorn, consists of a solution of gaseous ammonia in water. When freshly prepared the gas is freely given off, and serious results have followed its inhalation when large bottles containing the solution have been broken. The gas attacks the structures of the larynx and the mucous lining of the lower air-passages, often producing alarming dyspnoea, it is to this characteristic that the special features of ammonia poisoning, as compared with the effects produced by the fixed alkalies, are due. Monro and Workman<sup>1</sup> record three fatal cases of poisoning by gaseous ammonia which occurred in a cold storage from bursting of the pipe of a refrigerator through which ammonia gas was being forced by a pump. In two of the cases, death took place on the third day from asphyxia, caused by severe broncho-pneumonia. The dead body of the third man was found near to the fractured pipe.

**Symptoms.**—Immediately after swallowing strong ammonia water a violent burning sensation is experienced from the mouth down to the stomach, which is followed by vomiting, and often by purging, the ejected matter may contain blood. The epithelial layer of mucous membrane is at once stripped off and the mouth feels as though it were filled with “skins.” Some of the vapour is sure to be drawn into the larynx, producing a sensation of suffocation, which is followed by a real difficulty of respiration from tumefaction of the glottis, the breathing is noisy and stridulous, and the greatest distress is manifested from fear of impending suffocation. The voice is at once rendered feeble and hoarse, or it may be entirely gone. The patient is continuously occupied in attempts to rid himself of the detached membrane and viscid mucus which accumulate in the mouth. He sits up in bed with an extremely anxious expression, pointing towards the throat and intimating an earnest desire for relief. He suffers from intense thirst, which his condition disables him from alleviating. The symptoms of collapse are present—small pulse, cold, clammy surface,

<sup>1</sup> *Glasgow Med Journ.*, 1898.

sunken features, and, usually, great restlessness. If the case goes on to a fatal issue, coma usually supervenes, the breathing becomes more and more difficult, partly from swelling of the mucous lining of the air-passages, and partly from excessive secretion of mucus. Even when the symptoms are at their worst and death appears imminent, recovery may take place so far as the immediate condition is concerned, the breathing improves, the patient becomes able to swallow and to dislodge the mucus, and in a comparatively short time is out of danger. In one case of this kind, which was under the author's care in the Salford Royal Hospital, the patient (a woman), who had swallowed more than an ounce of commercial ammonia water, became gradually more and more collapsed, dyspnoëic, and cyanosed, until tracheotomy was contemplated. She recovered from the acute stage, but, later on, developed stricture of the œsophagus at its lower part. Another case ran a different and exceptional course. A woman aged forty-two drank an amount not exceeding one ounce of ordinary liquor ammoniæ the day before admission. Her chief distress was pain and difficulty in swallowing. For ten days she continuously improved, there was neither vomiting nor diarrhœa. On the eleventh day she suddenly became collapsed, hæmorrhage from the bowels occurred, but there was no hæmatemesis. She died in a few hours. On section the œsophagus was found to be extremely soft and friable at its lower end. The stomach was distended with blood clot, at the cardiac end the stomach wall was very thin and fragile. The duodenum was entirely filled with blood clot, but its walls were healthy. The rest of the bowel contained blood, the walls being also healthy. Pregnant women frequently abort in consequence of ammonia poisoning.

When recovery from the acute stage takes place, *apepsia*—due to destruction of some of the gastric glands—and stricture of the œsophagus or of the pylorus are to be feared.

**Fatal Dose.**—This is difficult to estimate, as the amount of gas present in a given quantity of the solution is subject to great variation. Two drachms have proved fatal, and recovery has taken place, as in the above-mentioned case, from more than one ounce. Death has taken place in a few minutes. The usual period for acute cases is twenty-four to forty-eight hours, when death is due to secondary effects life may be prolonged for years.

**Treatment** is the same as in the case of the fixed alkalis, special attention being paid to the respiratory symptoms, such as placing the patient in a tent, the air of which is rendered moist with steam.

**Post-mortem Appearances.**—If death takes place in the *acute* stage the lips are swollen, the mucous membrane of the mouth is softened and more or less detached, a similar condition existing along the œsophagus and, possibly, in the stomach. Immediately after the poison is taken the mucous membrane presents a white appearance, but it quickly changes to an angry-looking red, the whole thickness or the epithelial layer only of the membrane may be detached. In severe cases the muscular coat of the œsophagus or of the stomach is also softened and completely disintegrated. actual perforation is of exceptional occurrence. The effects of the poison rarely pass beyond the stomach. The laryngeal mucous membrane is infiltrated and thickened, it has been observed to be eroded and, in some cases, to be covered with exudation, forming a kind of false membrane. The condition of the laryngeal mucous membrane is usually attributed solely to the action of the vapour, but the appearances in one or two cases that the author had seen led him to believe that some of the liquid ammonia may not unfrequently be drawn into the larynx by spasmodic attempts at inspiration due to the extreme irritation set up by the vapour at the moment

of swallowing the poison. The smaller bronchi have been found to contain tubular casts. The kidneys may be inflamed. The **chronic** cases show corresponding post-mortem indications to those met with under similar conditions in poisoning with the fixed alkalies.

**Chemical Analysis.**—Ammonia is recognised by its odour, it may be separated from organic admixture by distillation, if the solution containing the ammonia is not alkaline, it may be neutralised with calcined magnesia before distilling. The gas is received into water acidulated with hydrochloric acid, and the ammonia is afterwards precipitated with platinic chloride in excess, the precipitate, being washed with alcohol to remove the excess, dried and weighed. 100 parts correspond to 15.68 parts of  $\text{NH}_4\text{OH}$ . The ammonia separated by distillation may be estimated volumetrically, if preferred. If putrefaction of the tissues is in progress it is of little use to examine them for ammonia as a poison, since it is evolved from decomposing nitrogenous organic matter.

**Tests.**—Ammonia responds like potash to platinic chloride, and to tartaric acid, it gives a brown precipitate with Nessler's reagent, and white fumes in the presence of gaseous hydrochloric acid.

**Ammonium Carbonate**  $[(\text{NH}_4)_2\text{CO}_3]$  when swallowed in large doses produces symptoms and organic changes resembling those produced by uncombined ammonia.

## CHAPTER XXX

### IRRITANTS.

#### SALTS OF POTASSIUM.

**Potassium Nitrate** ( $\text{KNO}_3$ ), saltpetre, or sal prunella, when swallowed in doses of one ounce or more, produces violent pain in the stomach and abdomen, with vomiting and purging, the ejected matter sometimes containing blood. Collapse occurs and is evinced by cold surface bedewed with sweat, and small, rapid, irregular pulse, which may subsequently become slow. Occasionally laboured respiration, unconsciousness and convulsions, pains in the loins, cramps in the calves of the legs, muscular twitchings, paræsthesiæ, paralysis of the limbs and aphonia have been observed. Death is usually preceded by coma, it may take place suddenly from heart paralysis. Gastric derangement may persist for a considerable time after the acute symptoms have subsided.

**Fatal Dose.**—The smallest recorded fatal dose is two drachms, which caused the death of a man aged forty. Recovery has taken place after one ounce. Death has occurred in from five to sixty hours.

**Treatment.**—Empty and wash out the stomach, give ice and opium to allay the sickness and pain, and alcohol if necessary. Mustard may be applied over the region of the stomach. Warmth and the recumbent posture are to be maintained.

**Chemical Analysis.**—If fluid, the suspected substance is filtered, if pul-taceous, it is extracted with water and then filtered. The filtrate is evaporated to a small volume and the salt allowed to crystallise out. The crystals are tested for nitric acid and for potash.

**Potassium Chlorate** ( $\text{KClO}_3$ ) possesses peculiar toxic properties. When taken in large doses, the red blood corpuscles are broken up by it, and the

hæmoglobin is converted into methæmoglobin. The mode in which the effects are produced is still in doubt, some observers hold that the salt is not decomposed within the organism, and, therefore, that the toxic effects are due to its specific action, others have demonstrated by experiments that when mixed with certain organic substances, such as pus and fibrin, the salt gives off oxygen. Binz<sup>1</sup> states that after prolonged action of these substances on potassium chlorate, the presence of chloric acid cannot be demonstrated. If blood is mixed with potassium or sodium chlorate to the amount of 4 per cent, it becomes syrupy, and on spectroscopic examination shows the bands of methæmoglobin, either alone or in combination with those of oxyhæmoglobin. The stroma of the red corpuscles parts with the hæmoglobin, which is subsequently changed into methæmoglobin, and the débris of the corpuscles produces certain pathological conditions which will be described among the post-mortem appearances.

**Symptoms.**—When large doses are taken the first symptoms are those of gastro-intestinal irritation, vomiting, pain in the stomach and bowels, with more or less collapse. Shortly after, pain is felt in the lumbar region, the urine, which may contain albumin, is diminished, also suppressed, hæmoglobin, methæmoglobin, and hæmatin have been found in it. The skin becomes cyanosed, and subsequently it is frequently jaundiced. Landerer<sup>2</sup> considers the jaundice to be partly polycholic and partly hæmatogenous. Palpation may reveal enlargement of the liver and the spleen. The patient is delirious, or apathetic and somnolent. At this stage the blood is brownish in colour and somewhat viscid, examined microscopically, colourless erythrocytes (the stroma of the red corpuscles) are seen along with normal corpuscles, interspersed with granular particles of free hæmoglobin, or of methæmoglobin. The white corpuscles are increased in number. The combined spectra of hæmoglobin and methæmoglobin will be present. Recovery may take place even when the symptoms have arrived at a very critical stage, when death occurs, an interval of several days usually elapses after the reception of the poison. Jacob<sup>3</sup> records the case of a woman aged thirty-nine who, after taking about twenty-five grammes (six and a half drachms) of potassium chlorate, became blue in the face, ears, fingers, and toes, she suffered from severe dyspnœa and the pulse was thready. At first active leucocytosis occurred, subsequently it almost ceased. The red corpuscles became paler and paler till scarcely a normally coloured one was present, "shadows" were numerous, granular masses of colouring-matter, both free and also within the stroma of the corpuscles, were seen, the red corpuscles which escaped granular changes showed poikilocytosis. On the fifth day the red corpuscles amounted to only 2,225,000, whilst the leucocytes reached 14,800 per cubic millimetre, the hæmoglobin was reduced to 20 per cent. At the first, methæmoglobin, but after the second day, only hæmoglobin, was present in the urine, which in red corpuscles were also seen. On the sixth day after taking the poison the patient died suddenly, and at the autopsy the spleen and kidneys were found enlarged and the lungs were gorged with blood. Ashby<sup>4</sup> saw a child fourteen months old that had been given about five grains of potassium chlorate three times a day for three weeks. It was anæmic, the gums were spongy and from them oozed brownish-coloured blood. The urine was brownish in colour and the liver and spleen were much enlarged. Death occurred from cardiac syncope. Schachtrupp<sup>5</sup> records two cases in which death followed

<sup>1</sup> *Arch f exp Pathol*, 1879

<sup>2</sup> *Deut Arch f klin Med*, 1891

<sup>3</sup> *Berliner klin Wochenschr*, 1897

<sup>4</sup> *Edin Med. Journ*, 1899

<sup>5</sup> *Dissert*, Halle, 1896

the use of potassium chlorate as a gargle without any being intentionally swallowed Rosselli<sup>1</sup> records a case in which the free use of a gargle of concentrated solution of potassium chlorate was followed by toxic amblyopia, recovery took place

**Fatal Dose** is uncertain, in one case six and a half drachms, and in another one ounce and a half respectively, caused death The fatal period ranges from six hours to as many days

**Treatment.**—The stomach should be emptied and washed out The after treatment will be symptomatic Diuretics and vapour baths, with fomentations or dry-cupping of the lumbar region, may be indicated

**Post-mortem Appearances.**—The mucous membrane of the **stomach** has been found swollen, softened, and easily separated from its bed, it may exhibit small ecchymoses The mucous membrane of the duodenum may present a similar swollen and softened appearance The **blood**, dark-brown in colour, is inspissated and sticky, when diluted with water it is chocolate-coloured and yields the spectroscopic reaction of methæmoglobin, under the microscope a large number of altered corpuscles are seen, which for the most part are shrunken and of knotty outline, together with numerous small free granules The **kidneys** are also chocolate-coloured, and on section show the most intense colour in the medullary portion, the glomeruli as small dots are visible to the unaided eye Examined microscopically, the straight and convoluted tubules are filled with a reddish-brown deposit formed of the debris of the red blood corpuscles, the epithelium is swollen and cloudy The **spleen** has been found enlarged and of a peculiar reddish-brown colour, besides normal red corpuscles the pulp may contain colourless erythrocytes The **liver** may be enlarged, its cells showing cloudy swelling with occasional pigmentation

**Chemical Analysis—Tests.**—The presence of potassium chlorate may be ascertained by adding a few drops of sulphuric acid to a solution containing the salt, and sufficient indigo sulphate to produce a moderately deep blue colour, two or three drops of sulphurous acid added to the mixture cause the colour to disappear If in organic admixture, some of the potassium chlorate may be separated by dialysis and examined as above

Several other salts of potassium and sodium—as the chlorides, sulphates, and carbonates—act as irritants when swallowed in large doses

## BARIUM.

**Barium Chloride** ( $\text{BaCl}_2 \cdot 2\text{H}_2\text{O}$ ) has been taken in mistake for Epsom salts and other saline purgatives It has been taken for suicidal purposes in the form of rat poison, into the composition of some varieties of which it enters

**Symptoms.**—In poisonous doses it acts locally as an irritant, and centrally as a nerve-poison In from a few minutes up to an hour or more after the poison is swallowed, violent pain is felt in the stomach and abdomen, accompanied by extreme nausea, which is followed by severe vomiting and purging The action of the heart is feeble and irregular, and pain may be experienced in the cardiac region The respirations are slow and laboured, the bronchi may become filled with mucus, causing dyspnoea and cyanosis The implication of the nervous system is further shown by ringing in the ears and diplopia, pains in the limbs, coma, convulsions, and, in some instances, paralysis Tactile sensation is often unaffected, and in the early stage the intelligence remains unimpaired

<sup>1</sup> *Bollet. dell'Ospedale oftalm. di Roma*, 1903.



Barium is eliminated by the kidneys and bowels, some of the amount taken is deposited in the bones Mendel and Sicher<sup>1</sup> state that barium is eliminated feebly by the kidneys, but more copiously by the bowels, though the rate of elimination is slow

**Fatal Dose.**—One teaspoonful of the powder has caused death A man aged fifty-five died in ten hours after swallowing a mouthful of a solution containing 130 grains of barium chloride (Stern)<sup>2</sup> The fatal period has been as short as one hour, and has extended to seven days

**Treatment.**—The stomach-tube should be used, or an emetic given unless vomiting has spontaneously occurred Sodium sulphate or magnesium sulphate should be given in large doses—half an ounce to an ounce or more Hypodermic injections of morphine and external warmth are useful

**Post-mortem Appearances.** The mucous membranes of the stomach and of the duodenum may be swollen and diffusely injected, or spotted with ecchymoses, in one instance the stomach was perforated, on the other hand death from a teaspoonful of barium nitrate has occurred in five hours without any significant hyperæmia of the gastric mucous membrane, or other indications of irritation, being found at the autopsy Particles of the poison have been found in the stomach in a case in which it was taken in the form of barium carbonate

**Chemical Analysis—Tests.**—The presence of a salt of barium in organic admixture can be readily ascertained by rolling up the end of a piece of thin platinum wire into a small ball, dipping it into the mixture, and transferring it to a Bunsen flame, a minute quantity of barium reveals itself by colouring the flame green, the experiment is best conducted in a darkened room If the green flame is examined with the spectroscope, the flame-spectrum of barium will be seen After taking up and drying some of the suspected fluid on the ball, the reaction will be rendered more distinct by dipping the wire in strong HCl

A solution of barium in organic admixture may be evaporated to dryness and incinerated, the product is drenched with  $\text{HNO}_3$ , and the excess of acid volatilised The nitrate, dissolved in water, may then be tested by adding dilute  $\text{H}_2\text{SO}_4$ , or an alkaline sulphate, either of which produces a white precipitate insoluble in  $\text{HNO}_3$ , it is also precipitated by a solution of KOH, a solution of potassium chromate produces a yellow precipitate insoluble in acetic, but soluble in hydrochloric acid If the barium is present as phosphate, or sulphate, it must be boiled with a concentrated solution of potassium carbonate for some time, and filtered, the filtrate is precipitated with dilute  $\text{H}_2\text{SO}_4$ , and the insoluble substance on the filter is dissolved in HCl, diluted with water, and also precipitated with  $\text{H}_2\text{SO}_4$  The combined precipitates, after washing and ignition, are weighed, this gives the amount of barium present as sulphate The absence of strontium and calcium should be ascertained by means of the spectroscope Death has occurred within thirteen hours after the victim had swallowed more than a teaspoonful of barium nitrate, and yet chemical examination of the viscera failed to reveal any trace of its presence

Other salts of barium—the carbonate, nitrate, and acetate—have acted as poisons

**Strontium** salts cannot be regarded as poisonous, the bromide, lactate, and nitrate have been administered medicinally in large doses, as much as 200 grains of the nitrate have been given in one day

<sup>1</sup> *Amer Journ of Physiol*, 1906

<sup>2</sup> *Zeitschr f Med Beamte*, 1896

## MAGNESIUM.

**Magnesium Sulphate** ( $\text{MgSO}_4 \cdot 7\text{H}_2\text{O}$ ), or Epsom salts, is usually regarded as a harmless purgative, but in large doses it has caused death. In experiments on animals it has been found to paralyse the respiratory movements, and also those of the heart. The former of these effects is well illustrated by the following case reported by Sang<sup>1</sup>—A woman, aged thirty five, dissolved four ounces of Epsom salts in some warm water and drank the solution. When seen shortly after she complained of a burning pain in the stomach and bowels, and a choking feeling, with a sensation as though she was losing power of her legs and arms, there was neither vomiting nor purging, the pulse was 96 per minute. An emetic of zinc sulphate was administered, but did not act, and before the stomach tube could be obtained profound collapse occurred. The pupils were dilated, the muscles of the face twitched, and there was complete paralysis, she then became comatose, and died in one hour and twenty minutes after swallowing the salts. The radial pulse was felt two or three minutes after the respirations had ceased.

Christison relates the case of a boy, aged ten, who, after swallowing two ounces of Epsom salts, was observed to stagger and appear very ill. Half an hour after, the pulse was scarcely perceptible, and the respirations were slow and laboured, in ten minutes more he died without vomiting having taken place. Luff<sup>2</sup> investigated the case of a girl aged twenty who apparently died from syncope in consequence of taking one ounce of Epsom salts on an empty stomach, the gastric mucous membrane was not inflamed.

## ARSENIC.

**Metallic Arsenic** is probably not poisonous, but, as it easily undergoes oxidation in the digestive tract, it may produce the usual symptoms of arsenical poisoning. A preparation known as fly-powder, which consists chiefly of finely divided metallic arsenic, probably along with some arsenious oxide, is strongly poisonous.

**Arsenious Oxide** ( $\text{As}_2\text{O}_3$ ), often called arsenious acid, or white arsenic, is the form in which arsenic is chiefly used as a poison. When fresh, it is a glassy-looking substance, with smooth vitreous fracture, after being kept some time it becomes white and opaque, resembling porcelain. In the form of powder it looks like flour, for which on several occasions it has been mistaken with fatal results. It has little taste, and having no colour, is easily administered for homicidal purposes, it has one compensating property—that of slight solubility in water. The solubility of arsenious oxide varies according to its molecular state, which is not constant, its density slightly diminishing with age, consequently, not only do the transparent and opaque varieties vary in respective solubility, but each individual specimen tends to differ from the rest. When the opaque variety is boiled with water for some time, 11.5 parts are dissolved per 100 parts of water, on cooling, from 2.5 to 3 parts of arsenious oxide per 100 of water are retained in solution, crystalline arsenious oxide is slightly less soluble. Cold water dissolves about  $\frac{1}{2}$  grain to 1 grain of arsenious oxide per ounce, the solution slightly reddens litmus paper. In acids and in alkaline solution arsenic is much more soluble. The law requires that, when arsenic is sold in quantities of less than 10 lbs, it shall be mixed either with soot or with indigo, at the rate of  $1\frac{1}{2}$  ounces respectively to the pound of arsenic.

When powdered arsenious oxide is added to water, or to liquid food, some of the finer particles float on the surface, forming a kind of white scum which cannot be got rid of by stirring, the appearance produced is very significant, and should be remembered when examining fluids which are suspected to have had arsenic added to them.

When combined with potash or soda, arsenious oxide is relatively much

<sup>1</sup> *The Lancet*, 1891.

<sup>2</sup> *Loc. cit.*

more soluble, and in this form it is used for domestic and other purposes, as well as in medicine. Some kinds of "fly-papers" are saturated with sodium or potassium arsenite, and much stronger solutions of arsenic can be obtained by soaking such papers in small quantities of water than by dissolving arsenious oxide itself. This fact has been utilised for criminal purposes.

**Arsenic Acid**  $[\text{AsO}(\text{OH})_2]$  is used in the manufacture of aniline colours, and, in combination with sodium, as a fly poison; it is less poisonous than arsenious acid.

**Arsenious Sulphide** ( $\text{As}_2\text{S}_3$ ), or orpiment, is almost insoluble in water, and when pure is said to be poisonous; the commercial variety usually contains uncombined arsenious oxide.

**Copper Arsenite** ( $\text{CuHAsO}_3$ ), or Scheele's green, and a mixture of copper arsenite and copper acetate, known as Schweinfurt green, though insoluble in water, are partially dissolved by the gastric juice. These pigments have produced toxic effects by their presence in the atmosphere in a state of minute division, the particles being derived from wall-paper or fabrics coloured with them. Copper arsenite dissolved in ammonia-water changes from green to blue, which is evidence of the presence of copper; on adding a crystal of silver nitrate, yellow silver arsenite forms round the crystal.

**Arsenuretted Hydrogen** ( $\text{AsH}_3$ ) is extremely toxic; it has produced poisonous effects in workmen who have been subjected to its influence when using hydrochloric acid contaminated with arsenic, to prepare iron for "galvanising," and also in the separation of silver from zinc. Serious results have followed its accidental inhalation in the laboratory.

**Sodium Cacodylate**  $[\text{As}(\text{CH}_3)_2\text{NaO}_2]$ , and some other organic compounds of arsenic, are but slightly toxic. Six to ten grains of sodium cacodylate (about equal to 4 to 6 grains of  $\text{As}_2\text{O}_3$ ) have been administered daily for several consecutive weeks without ill-effects. On the other hand, symptoms of arsenical poisoning have been caused by very much smaller doses. There are other organic compounds of arsenic which are virulent poisons.

Eczematous eruptions on the legs have been caused by wearing stockings coloured with aniline dyes prepared with arsenic.

**Salvarsan**, or dioxo-diaminoarsenobenzol, which is now extensively used in the treatment of syphilis, has been responsible for a considerable number of deaths, most of which are attributed by Mentberger<sup>1</sup> to arsenical poisoning.

Strathy, Smith and Hannah<sup>2</sup> have reported a series of 58 cases of poisoning by salvarsan, eight of which were fatal. Of the patients who died, one was under 20, four between 20 and 30, and three between 30 and 40 years of age. The greatest number of doses of salvarsan given was 11, and the least 4. The largest amount of salvarsan administered was 6.95 grammes, and the smallest 2.2 grammes. The average time onset of symptoms after the last dose was 41 days, the longest interval 48 days, and the shortest 18 days. The time which elapsed between the onset of the symptoms and death ranged from two to eleven days, the average being five days. The symptoms were similar in every case, sudden onset of jaundice being rapidly followed by nausea, epigastric pain, stupor, hæmatemesis, delirium, and death. Post-mortem, the most striking feature was acute atrophy of the liver and degenerative changes in its parenchyma.

In the 50 non-fatal cases, jaundice was the most prominent symptom, being exhibited by 39. It appeared within ten days of the onset of toxic symptoms, and lasted on an average for four weeks. Eight cases exhibited dermatitis,

<sup>1</sup> *Entwicklung und gegenwärtiger stand der Arsentherapie der Syphilis*, 1913.

<sup>2</sup> *Lancet*, April, 1920.

two showed peripheral neuritis, and two nephritis. The writers consider the symptoms to have been due to delayed arsenical poisoning.

### Acute Arsenical Poisoning.

**Symptoms** - The interval between the reception of the poison and the first appearance of the symptoms is determined by several conditions. In a state of solution the same dose acts more quickly than when administered in the solid form, the presence of food in the stomach tends to retard, and the absence of it to accelerate, the onset of the symptoms. The interval ranges from ten minutes, or even less, when a strong solution is swallowed on an empty stomach, to twelve or eighteen hours under converse conditions. If a poisonous dose of undissolved arsenious oxide is taken into a full stomach, and the recipient immediately after retires to bed and sleeps an unusually long period of quiescence may follow. The **usual period** is from **half an hour to an hour**. In a **typical case** a sensation of heat rapidly developing into a violent burning pain, is felt in the throat and stomach, then follows nausea, with uncontrollable vomiting and a sensation of constriction in the gullet. The vomited matter at first consists of food that may be present in the stomach along with opaque white masses consisting of mucus mixed with arsenious oxide, if the poison has been administered as a powder or only partially dissolved. If commercial arsenic mixed with soot or indigo-blue has been taken, the early vomit will probably be correspondingly tinged. After the contents of the stomach have been ejected, the vomit may consist of slimy mucus, or of fluid resembling rice-water, which may contain blood, or it may be bile-stained. Soon after the commencement of the vomiting, purging sets in, accompanied by distressing tenesmus, and, frequently, by a burning sensation in the rectum, after the evacuation of any *fæces* contained by the bowels, the dejections tend to assume the rice-water appearance, and they may contain blood. The pain in the stomach is usually, but not always, increased on pressure. The patient suffers from intense thirst attempts to relieve it being immediately followed by rejection of the swallowed fluid.

The feeling of sinking or depression, which precedes the primary vomiting, develops into one of extreme prostration and collapse, the face presents an appearance of great anxiety, the features are sunken, the surface is cold, moist, and cyanosed—especially the limbs—the pulse is small and thready, respiration is laboured, and the voice is hoarse. As a rule, the tongue is at first thickly coated with a white fur, subsequently it often becomes red at the tip and round the edges, sometimes it is unnaturally red over its entire surface. Owing to the profuse vomiting and purging and the reduced arterial tension the urine is scanty, and it may contain albumin or blood, attempts to urinate are painful. Cramps, especially in the calves, torment the sufferer, who tosses about to obtain relief. Death may be preceded by coma, which is not unfrequently accompanied by clonic or tonic spasms, or consciousness may be maintained to the end.

It will be observed that in many respects the symptoms strongly resemble those of cholera, when that disease is epidemic, errors in diagnosis may easily be made. If doubts arise in the medical attendant's mind, he should examine the excretions for arsenic.

The above description of the symptoms of acute arsenical poisoning embraces the principal features of a typical case, it is not to be inferred, however, that all these symptoms invariably occur in every case, nor that their progress is always precisely the same.

In exceptional instances the poison seems to spend its force on the **nerve-centres**, the gastro-enteric symptoms being less pronounced, or they may be entirely absent. In such cases, from the very first extreme collapse occurs with superficial and deep anæsthesia of the limbs, faintness, extremely small, feeble pulse, and coma, which quickly comes on, and terminates in death within six or eight up to twenty-four hours. Complete general paralysis may be present some hours before death.

In criminal poisoning **repeated doses** of arsenic may be given, a mode of administration which considerably modifies the course of the symptoms. The early doses produce gastro-enteric disturbance shown by vomiting, purging, pain in the stomach, foul tongue, loss of appetite, and a feeling of depression and languor, as the symptoms are passing off, but probably before they have quite subsided, another dose of the poison is given and the acute symptoms are renewed. In these cases, some of the symptoms of chronic arsenical poisoning are also present, there may be itching and smarting of the eyeballs and the margins of the lids, the conjunctivæ being reddened and granular, a similar hyperæmic condition of the mucous membrane of the fauces and throat causes constant hawking, apart from actual vomiting, the patient having a sensation as though a hair was in his throat. The tongue and mouth are dry, the former being either thickly coated or red and irritable looking. The skin has an unhealthy, half-jaundiced hue and may display erythematous or eczematous eruptions. More pronounced symptoms of neuritis occur in the acute form of poisoning—Creeping and tingling, especially in the fingers, numbness of the hands and feet, severe cramps, not limited to the calves, dropping of the ankles when the patient is lying on his back in bed, and extreme tenderness of the muscles on pressure. The patient is very restless and cannot sleep, and in the early stage the temperature will probably be slightly elevated.

The acute and chronic symptoms may be combined in variable proportion not necessarily corresponding to the length of time the patient has been under the influence of the poison. In some cases in which life has been prolonged for several days after the first inception of the poison, there may be almost entire absence of the chronic features, while in others several may appear within the first twelve hours. Diarrhœa, instead of coming on at once, may be delayed for one or more days, the abdomen being either tender or free from tenderness in the interval. In the case of *Reg v Maybrick* (Liverpool Assizes, 1889), in which the accused was found guilty of having caused the death of her husband by the administration of arsenic, evidence was given that purging did not occur until the third or fourth day, the abdominal pain being less violent than is usually the case. There was also absence of cramps in the calves, which, however, is a less constant symptom than diarrhœa. In the case of the Duc de Praslin,<sup>1</sup> the abdomen was painful and distended during the first four days after taking a fatal dose of arsenic, but there was only one evacuation of the bowels. Anderson<sup>2</sup> records a case where after nearly complete recovery had occurred death took place on the fifth day from intestinal hæmorrhage.

The combinations of arsenic with copper produce the usual symptoms of arsenical poisoning. In one case related by Seidel,<sup>3</sup> a girl aged nineteen swallowed a tablespoonful of mixed paint, the basis of which was Schweinfurt green, she died in sixteen hours, indications of the presence of the pigment were found in various parts of the digestive tract. Huber<sup>4</sup> relates the case of a man

<sup>1</sup> *Annales d'Hygiène*, 1847

<sup>2</sup> *Lancet*, 1910

<sup>3</sup> *Maschka's Handbuch*, Bd 2

<sup>4</sup> *Zeitschr. f. klin. Med.*, 1888

who took about 4 grammes of Schweinfurt green, he recovered from the immediate effects, but suffered severely from arsenical paralysis.

Death has resulted from the application of arsenical paste to destroy morbid growths, and in infants from the use of nursery-powder adulterated with powdered arsenious oxide. Arsenic has been used as a cosmetic, in the case of female poisoners with arsenic, any of the substance traced to their possession is generally accounted for on this ground. Applied as a solution for a limited time to the unbroken skin of an adult, it would not be absorbed to any dangerous extent, if at all. Very exceptionally arsenic has been surreptitiously introduced into the vagina with homicidal intent and has caused death. This mode of administration is of ancient origin, a tract printed in 1598 gives an account of the trial and condemnation of one Henry Robson, a fisherman of "Ryl," for thus poisoning his wife, who died five days afterwards. Haberd<sup>1</sup> records a unique case of a girl aged twenty-four who inserted arsenic into her vagina with suicidal intent and died two days after admission into hospital, isolated hæmorrhages and advanced fatty changes were found after death. Chabinat<sup>2</sup> records a case of fatal poisoning by arsenic from the application to the breast of an ointment composed of arsenious sulphide and butter. Arsenic was found in the internal organs. If the orpiment was pure the case proves that, although insoluble, it is not so inactive as is generally supposed.

**Fatal Dose.**—Two grains of arsenious oxide have proved fatal, recovery has taken place after a teaspoonful. When a fatal dose is taken the symptoms usually persist continuously until death takes place, which occurs in from twelve to forty-eight hours. Life is not unfrequently prolonged beyond the limit here stated, in such cases there are usually remissions in the course of the symptoms. The Duc de Praslin lived until the sixth day, Maybrick lived until the eighth day, and in exceptional cases death has not occurred until the fourteenth or even the sixteenth day. In one case, after a large dose, death took place in twenty minutes, it has not unfrequently occurred in from two to three hours.

**Treatment.**—Evacuate the stomach with the tube, or an emetic, and then give freshly-precipitated ferric oxide prepared by adding ammonia-water, or a solution of potassium carbonate, to the tincture of iron perchloride, the precipitate is strained off and administered suspended in water. Calomel and magnesia may be substituted if ferric oxide cannot be obtained. Demulcents and subsequently morphine should be given, together with the application of external warmth.

**Post-mortem Appearances.**—Externally the body may present a somewhat shrunken appearance, the eyeballs being sunk, and the surface may be slightly cyanosed, these appearances are by no means invariable. Rigor-mortis sometimes lasts unusually long. The important **internal signs** are afforded by the **stomach** and **intestines**. On opening the stomach, indications of intense inflammation present themselves, the whole of the mucous membrane may have a reddened velvety look, or the appearance may be limited to the greater curvature and the posterior part, or it may exist in two or more separate spots, the colour may be dark red or bright vermilion. Usually small dots or streaks of a darker colour are more or less numerous distributed over the inner surface of the stomach, which is frequently corrugated, this appearance is not invariable. It has been found absent in cases in which arsenic in a soluble form was administered. In some parts larger-sized submucous hæmorrhages may be seen. The surface of the mucous membrane is occasionally eroded, and particles

<sup>1</sup> *Wiener klin. Wochenschr.*, 1897.

<sup>2</sup> *Annales d'Hygiène*, 1890.

of undissolved arsenious oxide are not unfrequently found embedded at or near the spots so attacked. Very rarely, the inflammatory condition has gone on to gangrene or to perforation, softening of the mucous membrane, so that it can be easily separated from its bed, is less rare, and the wall of the stomach is sometimes thinned. The more profound lesions of the stomach wall are caused by the local action of the poison present in the solid form, in addition to the changes which are due to absorption. When a fatal dose of arsenic has entered the system by some other channel than the mouth, the post-mortem evidence of gastritis is nevertheless present, like some other poisons, arsenic is partially eliminated through the stomach, irrespective of the mode of its administration. The **duodenum** generally participates in the appearance presented by the stomach, the signs of inflammation may be limited to a few inches below the pylorus, or they may extend the whole length of the duodenum. The **jejunum** has also been found inflamed, and the rectum frequently so. Along with signs of diffuse inflammation in the intestines, small submucous hæmorrhages may sometimes be seen, the solitary glands and Peyer's patches are often swollen. The **œsophagus**, as a rule, is not inflamed. The **liver** and **kidneys** may yield microscopic evidence of granular or fatty degeneration, but such a condition is not usually distinguishable in rapidly fatal cases, in one case in which death took place in three hours Gumprecht<sup>1</sup> found desquamative papillary catarrh analogous to the "cholera kidney." None of the other organs show any characteristic changes.

In those exceptional cases in which paralysis of the nerve centres is substituted for the usual gastro-enteritis, the appearance presented by the stomach may be relatively trivial. Milford<sup>2</sup> describes the post-mortem appearance of the stomach in a case of this kind, in which no vomiting had occurred, although after death the stomach contained not less than 200 grains of arsenic. Only about a quarter of the surface of the mucous membrane near the pylorus presented a bright scarlet colour, the remainder was normal, the duodenum presented a similar but less marked discoloration, the rest of the digestive tract was unchanged.

St George<sup>3</sup> reports the following remarkable case —

At 8 p.m. on February 23rd, 1920, a man aged 68, who carried on a pharmacy business, took by mistake for a preparation of magnesia a heaped teaspoonful of arsenic, presumably the trioxide, mixed with hot milk. Half an hour later he ate a hearty supper of porridge and milk. At midnight he had vomiting, diarrhœa, and a burning pain in the stomach. At 4 a.m., when seen by St George, his pulse was 120 and thready, his temperature was subnormal, and he had cramps in the legs. It was considered to be too late to wash out the stomach, but the symptoms were relieved by an enema containing tincture of opium, and he was given albumen water by the mouth. There was only one motion of the bowels, and this was not until late in the evening. The urine was suppressed. Diuretics were ordered, and magnesium sulphate was given freely. Urine was passed and the symptoms improved. The patient was soon able to get up and walk down to the doctor's house. At a later date he began to complain of pain in his arms and numbness of the hands, and was unable to hold a cup firmly, but he was still able to walk. Soon his feet felt as if he were walking on wool, and he was removed to a nursing home on April 30th. Shortly after admission he became ataxic, and was unable to stand upright or walk with his eyes closed. The knee-jerks gradually disappeared, and ankle clonus developed. He became slowly but steadily worse, hypostatic pneumonia developed, and death took place on June 13th by slowly ascending paralysis. Up to the end arsenic could be found in the urine.

**Arsenuretted Hydrogen** is a powerful blood poison which disintegrates the red blood corpuscles and sets free the hæmoglobin. The **symptoms** may come on immediately, or they may be delayed eight or ten hours after the gas has been inhaled, they comprise a feeling

<sup>1</sup> *Deutsche med. Wochenschr.*, 1893.

<sup>2</sup> *Australasian Med. Gaz.*, 1890.

<sup>3</sup> *Brit. Med. Journ.*, February, 1921.

of malaise, headache, and dizziness, shivering or prolonged rigors, vomiting, pain in the loins and epigastrium, the presence of blood colouring matter in the urine, jaundice, and usually constipation. The vomit and the motions often contain blood, the urine usually contains bile pigment, fatty casts, and free fat granules. The red blood corpuscles are greatly reduced in number, in one case there were only 1,800,000, and in another 920,000 per cubic millimetre. As  $\text{AsH}_3$  acts feebly, if at all, on the hæmoglobin it sets free, the jaundice is probably due to blocking of the bile ducts by the inspissated bile which is formed from the free hæmoglobin present in the blood plasma. **Post-mortem**, the mucous membrane of the stomach and small intestines is deeply congested and shows petechial hemorrhages, the liver and kidneys are swollen, and the viscera generally are superficially tinged blue or blue black, the lungs are cedematous. Microscopical examination shows a tendency to fatty changes, and chemical analysis reveals the presence of arsenic in the tissues. For a detailed account of arsenuretted hydrogen poisoning see a paper by the author and J. Gray Clegg in *The Medical Chronicle* for 1895.

### Chronic Arsenical Poisoning.

Arsenic may be received into the system in small quantities for a prolonged period, and thus produce symptoms differing from the acute form of poisoning. The sources from which the poison is derived comprise wall-papers, fabrics, artificial flowers, toys, and fancy papers which have been used to envelope confectionery. Trade risks constitute another source, as, for example, in the manufacture of arsenical sheep-dipping, which consists of crude sodium arsenite mixed with sulphur. Copper arsenite, either alone or in admixture with copper acetate, is the form in which the poison is used for colouring purposes, in the case of wall-papers and fabrics the pigment is usually so loosely attached that particles are freely given off, and, floating in the atmosphere, are inhaled and swallowed. Some years ago many cases of arsenical poisoning were thus caused, and public attention being drawn to the subject, manufacturers ceased to employ the dangerous pigment. The evil, however, is not altogether a thing of the past, Harding<sup>1</sup> records a number of cases of chronic arsenical poisoning which occurred among the nurses in an asylum, the symptoms being eventually traced to the use of some green baize curtains, which were found to contain a large amount of arsenic. Kuttner<sup>2</sup> reports a series of cases resulting from the use of arsenic in bedroom carpets. Certain moulds, the most active being *Penicillium brevicaulis* and *Mucor mucedo*, possess the property of developing volatile combinations with arsenic, and it has been suggested that chronic poisoning from arsenical wall-papers may be due to the inhalation of such products. This may occur, but the simpler explanation—inhilation of minute particles of arsenic—is more probable. In 1900 a widespread outbreak of peripheral neuritis associated with various affections of the skin occurred in Lancashire. Reynolds,<sup>3</sup> who was the first to recognise the cause, and to publish an account of the outbreak, rightly attributed it to arsenic, the presence of which he detected in various samples of beer. It was subsequently proved that the beers thus contaminated had been brewed from glucose and invert sugar manufactured by a firm that, in its preparation, had used sulphuric acid largely contaminated with arsenic. Numerous analyses showed that the contaminated beers contained from 3 grains to less than  $\frac{1}{100}$  grain per gallon of arsenious oxide. Luff<sup>4</sup> and others found that certain malts contained arsenic derived from the coke and anthracite coal used in drying them. Taylor and Trubshaw<sup>5</sup> record six cases of poisoning among girls working in a laundry which was heated by a coke stove.

<sup>1</sup> *The Lancet*, 1892.

<sup>2</sup> *Berlin klin. Wochenschr.*, 1912.

<sup>3</sup> *Brit. Med. Journ.*, 1900.

<sup>4</sup> *Royal Com. on Arsen. Poisoning*, 1901.

<sup>5</sup> *Brit. Med. Journ.*, 1911.



**Symptoms.**—The early indications consist of gastric disorders, loss of appetite, headache, general feeling of malaise, and constipation or diarrhœa, then follow colicky pains, irritation of the eyelids, cachectic hue of the skin, and eczematous eruptions, especially in the folds of the axilla, or between the scrotum and thighs, subsequently the skin becomes pigmented. Sooner or later pronounced indications of peripheral neuritis develop, the characteristic features of which are sensory disturbances, motor paralysis, and ataxia, the effects of arsenic on the nerves resemble those of alcohol, and differ from those produced by lead, in the prominence of the sensory disorders. The symptoms come on at variable intervals, from a week to three or four weeks, after the initial effects of the poison have manifested themselves, they usually commence with sensory disturbances—tingling, numbness, formication, and, in some cases, cutaneous anæsthesia. Then the affected muscles rapidly atrophy, the knee-jerk is usually lost, and the reactions of degeneration are present, according to G. Brouardel,<sup>1</sup> the legs alone are attacked in about 50 per cent of cases, in the arms the paralysis has the same distribution as in lead paralysis. The aggregate symptoms in the Lancashire outbreak were as follows—In a few instances, there was a history of gastro-intestinal irritation. An early symptom was suffusion of the eyes and the conjunctivæ, and the mucous membrane of the throat was reddened. Patients complained of a burning sensation in the hands and feet accompanied by tingling and numbness, and the palms of the hands and the soles of the feet were bright pink in colour and were moist with sweat (erythromelalgia). Erythematous, urticarial, and herpetic eruptions were common, occasionally a pemphigoid eruption appeared. Subsequently the horny layer of the epiderm thickened, especially on the palms of the hands and the soles of the feet, and then flaked off in large scales (hyperkeratosis), this was a very common symptom. Pigmentation, varying from a few freckles, or a slight darkening of the skin about the axilla and the groins, to a mulatto-like appearance, was also common. A full description of the skin symptoms is given by Brooke and Roberts.<sup>2</sup> The neuritis was characterised by its severity both as regards sensory and motor symptoms. The muscles, especially of the legs, were exquisitely tender, the weight of the bed-clothes being often insupportable. In the worst cases, the patients lay in a crouching attitude, with the knees flexed and drawn up. Recovery was very slow. In fatal cases, death frequently occurred with unexpected rapidity, from heart-failure.

When the action of the poison is continued the cachexia becomes more pronounced, anæmia, falling off of the hair, defective nutrition of the finger nails causing their detachment, and other trophic disturbances manifest themselves. The skin pigmentation is supposed to be due to deposition in the rete malpighii and in the lymphatics of the corium, especially about the papillæ, of granules of altered blood colouring matter, a certain proportion of hæmoglobin being converted by the arsenic into a substance, closely resembling bilirubin, of which the granules consist. On the other hand, Delpine<sup>3</sup> holds that a pigment—melanin—which is not derived from hæmoglobin, is elaborated as a physiological product in the deeper layers of the epidermis, though under normal conditions not to such an extent as to be visible, and that arsenic, by stimulating the epidermis, determines an excessive production of melanin with consequent pigmentation. After death Ehrlich and Rybalkin<sup>4</sup> found pigmentary changes in the ganglion cells of the anterior cornua in the cervical and lumbar enlargements, as well as degeneration in the peripheral nerves, in addition, Henschen<sup>5</sup> found the ganglion cells to be atrophied and even to have disappeared.

The strong affinity between keratin tissues and arsenic (see Elimination) suggested a possible explanation of the cause of the neuritis in chronic arsenical poisoning. The axis

<sup>1</sup> *Etude sur l'Arcenicisme*, 1897

<sup>2</sup> *Brit Journ of Dermatology*, 1901

<sup>3</sup> *Upsala Lakaref, Fordhandl*, 1893

<sup>4</sup> *Proc of the Physiol Soc*, 1890

<sup>5</sup> *Neurologisches Centralbl*, 1892

cylinder and the white substance of Schwann are covered with a sheath of neurokeratin and are also traversed by numerous transverse fibrils of this substance, which is closely allied to keratin. Brain substance likewise contains neurokeratin, and the fact that the white matter contains about ten times more than the grey matter was made use of to ascertain whether or not the same affinity exists between neurokeratin and arsenic as between keratin and arsenic. The brains from a number of cases of chronic arsenical poisoning were severally divided into two portions, one consisting chiefly of white matter and the other of grey. Equal weights of the two portions of each brain were separately analysed, with the result that the white matter invariably yielded more arsenic than the grey. For example 240 grms of white matter from one of these brains contained as much as 0.0008 gm of arsenious oxide, whereas 240 grms of the grey matter from the same brain yielded merely an unweighable amount. In brains which contained very little arsenic the difference was not so great. It thus appears probable that neurokeratin has a strong affinity for arsenic, this may exercise a considerable influence in the production of both brain and nerve symptoms in chronic arsenical poisoning.

Arsenic appears to interfere with tissue oxidation and thus to determine the occurrence of fatty changes. Binz and Schulz<sup>1</sup> hold that arsenic has the power of taking from and giving oxygen to the tissues. Binz<sup>2</sup> states that arsenious oxide is mostly excreted as arsenic acid. This is opposed by Husemann<sup>3</sup> and others, the author's investigations led him to the conclusion that most of the ingested arsenic is excreted as arsenious oxide. It is probable, as suggested by Mott with regard to phosphorus, that arsenic interferes with the power of cells to take up oxygen and to store it in their protoplasm, hence the fatty changes.

Typical multiple neuritis has been known to occur after a single poisonous dose of arsenic. Jolly<sup>4</sup> reports a case in which a woman aged 27, five days after swallowing a cupful of Schweinfurt green mixed with coffee, which produced the usual gastro enteric symptoms, began to have numbness of the feet, paresthesiæ of both hands and feet, followed by motor paralysis, ataxia, and marked atrophy of the muscles of the calves, subsequently her hair came off, and she had excessive sweating of the palms of the hands, recovery ultimately took place. Meirowitz<sup>5</sup> relates a case in which, three weeks after a man aged 19 swallowed a teaspoonful of powdered arsenious oxide, marked symptoms of peripheral neuritis developed, followed by considerable atrophy of the muscles of the legs and hands.

Nutt, Beattie, and Pye Smith<sup>6</sup> have collected 30 cases of cancer of the skin following the prolonged taking of arsenic. In nearly all of these hyperkeratosis, especially of the palms and soles, was present, in half the cases the cancerous lesions were multiple, and in a fourth of the cases the age of the patient did not exceed 35 years, again suggesting a special cause. Sir Jonathan Hutchinson referred to arsenic as a cause of cancer so far back as 1887.

**Arsenic-eating**—In relation to the chronic ingestion of arsenic, mention must be made of the alleged tolerance to the poison that may be acquired by its habitual use. It is stated that the peasants in Styria, by progressively augmenting the dose, acquire the capacity of swallowing with impunity four or five grains of arsenious oxide at a time, the object being to enable the "arsenic eater" to endure greater fatigue in mountain climbing than he otherwise could do. It is said that arsenic is sometimes given to horses in order to improve their coats and general appearance. Although the Styrian habit was authenticated by competent and trustworthy observers such as Professors MacLagan<sup>7</sup> and Roscoe,<sup>8</sup> the statements were so irreconcilable with the well known action of arsenic on the organism as to be received with much scepticism. Recently, however, Cloetta<sup>9</sup> has experimentally investigated the subject, and has arrived at a solution of the mystery. He found that arsenious oxide dissolved in water could not be administered to a dog in larger doses than 0.0035 gm without giving rise to toxic effects, but by giving the arsenious oxide in the solid form, at first in small doses and then in progressively larger doses, a daily dose of two grammes (31 grains) could, in about two years, be taken with impunity. By simultaneous analyses of the urine and fæces, Cloetta discovered the reason of the immunity as the dose of solid arsenic was augmented, its excretion in the urine diminished with a parallel increase in fæces. The immunity, therefore, is local, and is due to lessened capacity of the digestive tract to absorb the arsenic it receives. This was further demonstrated by injecting in solution one sixtieth of the accustomed daily dose, when the animal quickly died, moreover the viscera contained but a small amount of arsenic.

<sup>1</sup> *Arch f exp. Pathol*, 1879

<sup>2</sup> *Ibid*, 1897 and 1898.

<sup>3</sup> *Deutsche med Wochenschr*, 1892

<sup>4</sup> *Ibid*, 1893

<sup>5</sup> *The Journal of Nervous and Mental Disease*, 1895

<sup>6</sup> *Lancet*, 1913

<sup>7</sup> *Edin Med Journ*, 1864

<sup>8</sup> *Mem Lit and Phil Soc*, Manchester, 1860

<sup>9</sup> *Arch f exp Pathol*, 1906

The **elimination** of arsenic takes place by several channels, of which two—the **kidneys** and the **bowels**—rapidly come into action. Arsenic may also be found in the sweat, the saliva and bronchial secretion, and, during lactation, in the milk. Subsequently it appears in the skin and its appendages the **nails** and the **hair**. Arsenic may readily be detected in the urine half an hour after a single five-drop dose of liquor arsenicalis, equal to about one twenty-first of a grain of arsenious oxide, has been swallowed, after a like dose it may also be detected in the **fæces**. Arsenic has been regarded as a non-cumulative poison, and when only a few doses are taken, this view is correct. In cases of acute arsenical poisoning it is very exceptional to find arsenic in the urine longer than a week or ten days after the administration of the poison, and in those fatal cases in which the patient survived more than ten days arsenic has rarely been found in the tissues. “Cumulative,” however, is a relative term, and although arsenic does not appear to combine intimately with the tissues as some of the heavy metals do, still when small doses are taken into the system in close succession for a considerable period, as in chronic poisoning a certain residuum remains in the tissues, after the last dose has been taken, for a much longer time than was formerly supposed. In cases of arsenical beer-poisoning the author found arsenic in the urine as long as thirty-one days, and in one instance fifty-nine days after the reception of the last dose. The longest period after which arsenic was found in the tissues was in the case of a woman who survived fifty-two days after the last dose, in several instances the author found it after from fourteen to twenty-seven days’ survival. Since the elimination of arsenic begins very promptly, and is usually continuous, it is probable that prolonged retention of the poison in the system is due to low arterial tension, caused to some extent by the action of the poison on the heart and to coincident depressed molecular activity in the tissues. In chronic arsenical poisoning arsenic may be detected, over prolonged periods, in the exfoliated horny scales forty-nine days after the last dose, and in the nails and hair four months after. There is a strong affinity between the keratin-tissues and arsenic, well-defined crystals of arsenious oxide were obtained from 0.2 gm. of hair, 0.03 gm. (half a grain) of horny scales, and from 0.03 gm. (half a grain) of nail-clippings. The true skin retains relatively little arsenic, in a case in which there was pronounced hyperkeratosis the true skin contained an unweighable amount, whereas the horny scales yielded equal to 0.013 per cent and the hair equal to 0.008 per cent of arsenious oxide. The nails probably contained still more, judging from the deposits yielded by the small available weights of nail-clippings. The progress of elimination may be followed in the hair, at a certain period arsenic could be detected in 0.3 gm. from the ends of the long hair of a woman, whilst a like weight taken from close to the scalp afforded no trace. Arsenic is probably loosely held by the tissues. 0.1 gm. of horny scales was boiled for one minute in a few cubic centimetres of distilled water, which was then passed through close-textured filter-paper in order to keep back all solid particles, yet arsenic was found in the filtrate.

As regards the viscera, arsenic that has been absorbed is retained in the largest amount by the liver, in chronic as well as in acute poisoning, of the abdominal viscera the kidneys come next, and then the spleen. In chronic poisoning it is to be found in the brain and in spongy bone, such as the bodies of the vertebræ and the base of the skull, but neither in brain-substance nor in bone has the author found arsenic to be exclusively, or even prominently, present, as is stated by some observers, when only a minimum amount is

retained in the body, this is corroborated by the experience of Stevenson<sup>1</sup> It is noteworthy that the red marrow of bones is enormously increased in chronic arsenical poisoning

Gautier<sup>2</sup> states that arsenic is normally present, in weighable amount, in the thyroid gland of man and animals and, in lesser amount, in the thymus and the brain, he also states that it is absent from the greater part of the tissues and from the blood The first clause of this statement is not generally accepted It is possible that traces of arsenic may occasionally be found in some of the tissues without there being any obvious explanation of its presence, but this does not constitute arsenic a physiological component of the human body Until more evidence is forthcoming, arsenic must, as hitherto, still be regarded as not being a normal constituent of the body, and consequently, when detected in criminal cases, its presence has to be accounted for, moreover Gautier does not claim to have found arsenic to be normally present in the liver On the other hand, Kunkel<sup>3</sup> failed to find arsenic as a normal constituent in any animal organs, including the thyroid gland

When arsenic is found in disinterred bodies it has been suggested that its presence may be due to transudation of the poison from the soil which surrounded the coffin Arsenic has been found in the soil of certain graveyards, usually, however, in combination with iron and in insoluble form, it is, therefore, in the highest degree improbable that a body free from arsenic when buried should become contaminated from the soil of a cemetery or churchyard, but to avoid possibility of error in this respect, a sample of the soil which surrounds the coffin should be collected and examined

**Post-mortem Imbibition of Poisons**—Some poisons when present in the stomach of a dead body tend progressively to diffuse themselves through the walls of the stomach to the neighbouring organs, this is especially the case with arsenic The extremely improbable allegation—which, however, has been made (Peckham)<sup>4</sup>—that the arsenic found in the viscera of a supposed victim of homicidal poisoning was introduced into the stomach after death, has led to the performance of experiments on the dead bodies of infants, and of some of the lower animals, in order to compare the results of *intra vitam* absorption of poisons with those of *post mortem* diffusion The order in which organs become impregnated by transudation of poison, which has been introduced into the stomach after death, is determined by anatomical conditions the viscera of the left side become infiltrated before those of the right side, so that within certain time limits it may be possible, in a given case, to arrive at a conclusion as to whether the poison was introduced before or after death Thus Orfila observed that by post mortem imbibition the left lung is affected before the right Strassmann<sup>5</sup> found that in not later than twelve days after the poison was introduced into the stomach of a dead body the left kidney contained arsenic whilst the right remained free, the left lobe of the liver also became impregnated before the right lobe, the same results were obtained when the bodies experimented with were laid on the right side Strassmann doubts the possibility of post mortem transudation of arsenic from the stomach to the brain, at any rate he never found it to occur, although some of his experiments lasted four weeks, on the other hand, some observers state that it may reach the brain in from twenty five to thirty days If within the first few weeks after death the left kidney contains arsenic and the right contains none, the inference to be drawn is that the poison was introduced into the stomach after death, the same applies to the lungs, but here *post mortem* transudation of the poison from the throat, œsophagus, or the air passages may lead to impregnation of both lungs In order that the inference shall hold good the difference must be absolute—not relative the left kidney must contain arsenic and the right none, it is not enough that the right kidney shall contain less than the left, such a difference is compatible with vital absorption The most important differential sign distinguishing *ante-mortem* from *post mortem* ingestion of arsenic is to be looked for in the stomach and duodenum

<sup>1</sup> *Royal Com. on Arsen. Poisoning*, 1901

<sup>2</sup> *Zeitschr. f. Physiol. Chem.*, 1905

<sup>3</sup> *Comptes Rendus*, 1899 et 1900

<sup>4</sup> *Boston Med. and Surg. Journ.*, 1880

<sup>5</sup> *Virchow's Arch.*, 1894

in acute arsenical poisoning, the mucous membrane of these viscera will present the appearances described on p 374, if the poison was introduced into the stomach after death there will be no such appearances, because, being the result of vital processes, they cannot be produced in the dead body

The body of a person who has died from arsenical poisoning is accredited with the power of resisting putrefaction. If at the time of death the tissues contain much arsenic, it will exercise a preservative influence, but it is not to be assumed that decomposition is retarded in every case of arsenical poisoning, much of the poison would not be present in the body after a minimum lethal dose, and in such a case putrefaction would follow the ordinary course. Instances of delayed putrefaction in the bodies of those who have died from arsenic have frequently been observed. In the case of *Reg v Cross* (Munster Assize, 1887), for the murder of his wife by poisoning her with arsenic, Pearson<sup>1</sup> states that when the body was exhumed, seven weeks after death, all the organs were in an excellent state of preservation, the stomach and the intestines appearing as fresh as though the deceased had died but twenty-four hours previously, in this case the fatal illness lasted about three weeks, during which time repeated doses of arsenic had probably been given, chemical analysis showed that the body contained a large amount. Brouardel and Pouchet<sup>2</sup> examined the body of a woman who died from arsenical poisoning in May, the body was exhumed on the thirtieth of October following, and was found to be in a remarkable state of preservation, not a trace of the gases of putrefaction being present, arsenic had been administered for six weeks before death, and a considerable amount of the poison was found in the body.

**Chemical Analysis.**—If arsenious oxide has been administered in the solid form it is probable that undissolved particles may be found lying on, or embedded in, the mucous membrane of the stomach, if the body has been interred, it is possible that the oxide may have been converted into the sulphide. Partial conversion into sulphide may occur before burial. Letheby<sup>3</sup> found it thirty-six hours after death, Taylor<sup>4</sup> in twenty-one hours after death, Paterson<sup>5</sup> in twenty-four hours, Harvey Littlejohn<sup>6</sup> the second day after death, and in another case fifty-two hours after. Any such particles should be picked out or scraped off, dried and tested, particles of colouring-matter—as soot or indigo—with which the arsenic may have been mixed should be sought for.

**Tests.**—In the first instance, **Reinsch's method** of testing for arsenic may be adopted, by this test 1 1,000,000 and, by concentration of the solution, 1 7,000 000 may readily be detected. Urine should be evaporated down to one-fourth or one-sixth its volume before being tested, solid substances should be pulvified and mixed with water so as to render them fluid. Hair should be cut into minute fragments, and horny scales of epiderm and nail-clippings should be finely divided and distributed in a sufficiency of distilled water. Previous to testing the suspected substance, the reagents themselves *should invariably be tested for arsenic*. Pure copper is easily obtained but hydrochloric acid absolutely free from arsenic is the exception. To some water in a flask, one-sixth its volume of strong hydrochloric acid is added together with two small pieces of copper foil, the flask is placed on a support covered with wire gauze over a Bunsen flame, and the acidulated water is allowed to boil gently for half an hour. The copper is then examined. If it retains its primitive brightness and colour, absence of arsenic in the reagents may be assumed. The

<sup>1</sup> *Dublin Journ Med Science*, 1888

<sup>2</sup> *Annales d'Hygiene*, 1888

<sup>3</sup> *The Lancet*, 1847

<sup>4</sup> *Guy's Hosp Reports*, 1851

<sup>5</sup> *Edin Med Journ*, 1858

<sup>6</sup> *Ibid*, 1906

acidulated water is now replaced by some of the suspected fluid, to which one-sixth its volume of strong hydrochloric acid is added from the same bottle as that from which the previous supply was taken, one or both of the same pieces of copper being dropped into the flask, which is again placed over the flame. With some substances—notably with hair—violent “bumping” occurs which necessitates the flask being supported by a clamp, else it will be projected off the tripod. If much organic matter is present, especially in the case of urine that has been evaporated down, it will probably be advisable to increase slightly the proportion of acid. When the amount of arsenic is small only one piece of copper should be used. After gently boiling for from half an hour to an hour, the foil is again examined, if the amount of arsenic in the fluid was very small,

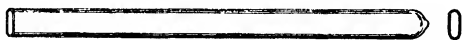


Fig 22—Sublimation tube

the foil will only exhibit a purple tint, if a little more arsenic was present, the foil will present a steel-grey appearance, if a large amount was present, it may be covered with a black amorphous coat which is easily detached. The foil is successively washed in distilled water, alcohol, and ether, is carefully dried on filter-paper, and then introduced into a small dry sublimation-tube, the form shown in the illustration is convenient for subsequent microscopical examination, the flattened walls producing less distortion than those of a cylindrical tube.

The closed end of the tube, on which the foil rests, is brought into the margin of a Bunsen flame and there retained until the film of arsenic is volatilised. When the arsenic leaves the copper, it combines with some of the oxygen of the air, and is deposited, as a ring—a centimetre or more up the tube, in accordance



Fig 23—Crystals of  $As_2O_3$

with the heat brought to bear—of octahedral or tetrahedral crystals of arsenious oxide. On microscopical examination the largest crystals will be found nearest the foil, where (unless the amount of arsenic is very small) the ring is sharply defined, the crystals, unlike those deposited from aqueous solutions, are always separate and distinct. A crystalline deposit, obtained as described is very characteristic of arsenic. Having removed the pieces of copper, a couple of drops of water are introduced into the tube and, with the aid of heat, the crystals are dissolved, this takes a few minutes to accomplish on account of the feeble solubility of arsenic, especially when in

the crystalline form. When all the deposit is dissolved, the solution is shaken out on to a colour-slab, so as to form two separate drops, to one a drop of a solution of silver nitrate is added, and to the other a drop of a weak solution of copper sulphate, a glass rod that has been dipped in ammonia-water is then held horizontally over the drops, close to (so as to allow the gaseous ammonia to act upon them), but without touching them. The one to which silver nitrate was added turns yellow, and the other becomes first blue and subsequently green, the salts formed are respectively arsenite and copper arsenite or Scheele's green. If a considerable amount of arsenious oxide is present, the drop to which silver nitrate is added will turn yellow immediately, before the application of the ammonia.

In addition to arsenic—antimony, mercury, silver, bismuth, platinum, palladium, tin, and gold are deposited on copper when boiled with it in acid solution; of these only three—arsenic, antimony, and mercury—yield sublimate: crystalline with arsenic, amorphous with antimony, and globular with mercury. Sulphur-yielding bodies in organic substances may cause the copper to be stained, therefore discoloration alone is never to be accepted as proof of the presence of arsenic. When arsenic is present as arsenic acid, it is extremely difficult to obtain any deposit on the copper. Reinsch's test is inapplicable in the presence of chlorates and nitrates. Attempts have been made from time to time to use Reinsch's test quantitatively. The insuperable difficulties are: that the copper foil does not abstract all the arsenic from the liquid, nor does it invariably give it all up in the sublimation-tube. The best quantitative results with Reinsch are obtained by Delépine's<sup>1</sup> method of comparing the sublimate obtained with others prepared under like conditions from known amounts of arsenious oxide.

**Marsh's test** is founded on the power possessed by nascent hydrogen of reducing arsenious and arsenic acids, and of combining with the liberated

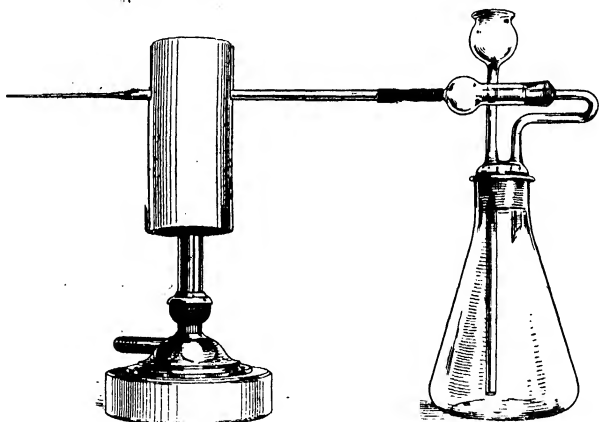


Fig. 24.—Marsh apparatus.

arsenic, forming arsenuretted hydrogen, from which the arsenic can afterwards be dissociated by heat and by chemical reagents. Arsenuretted hydrogen has a penetrating garlic-like odour, when it escapes into the air, which reveals its presence in the most minute traces. Marsh's test will detect the two-hundredth of a milligramme, or, according to some, the one-thousandth of a milligramme of arsenic. The necessary apparatus consists of a bottle or flask, through the stopper of which a long thistle-funnel and an exit-tube pass; between the flask and the free end of the exit-tube, which for the last two inches is reduced to a small diameter, a drying-tube filled with granules of chloride of calcium and plugged at each end with cotton-wool is interposed to dry the gas as it escapes. Some chemists recommend the further interposition of a tube

<sup>1</sup> *Brit. Med. Journ.*, 1901.

containing dried lead acetate paper, or cotton-wool moistened with a solution of lead acetate between the flask and the chloride of calcium tube in order to intercept sulphur and selenium compounds. The accompanying figure is taken from a convenient model which the author had constructed entirely of glass, in order to avoid the possibility of contamination from the accidental introduction of arsenic by the use of rubber stoppers. In 1861, Bloxam<sup>1</sup> published a volatile method of developing arsenuretted hydrogen for analytical purposes, and later W. Thompson<sup>2</sup> devised an electrolytic Marsh-apparatus, which is capable of yielding very delicate reactions. As the apparatus is somewhat complicated and is relatively costly, it is doubtful whether the electrolytic adaptation— notwithstanding the advantages which it possesses—will displace the ordinary Marsh-Berzelius method, there being little to choose between them as regards delicacy.

Three or four grammes of metallic zinc free from arsenic are placed in the flask, and some dilute sulphuric acid (1-8), also free from arsenic, is added, hydrochloric acid (1-3) is preferred by some. The purer the zinc, the less freely is it attacked by the acid, with some specimens it may be necessary to promote action by the addition of a single drop of a solution of platonic chloride to the contents of the flask, but rapid evolution of hydrogen being disadvantageous, it is better to allow the acid to act unaided if it will do so. In any case platonic chloride must not be added after the suspected fluid has been introduced into the flask, else some of the arsenic will be kept back. When all the air is expelled from the flask, a lighted Bunsen burner is placed under the exit-tube, so as to render a portion of it incandescent at about an inch from the narrowed part, the tube being supported on both sides of, and near to, the flame, but if a trace of air is present in the flask when the flame is applied to the tube, water will be formed in it to the detriment of the arsenical deposit. In order to prove the absence of arsenic in the acid and the zinc, the flame is allowed to play on the tube for at least thirty minutes before any of the suspected substance is introduced into the flask, if the tube remains free from deposit, the purity of the reagents is established. A little of the suspected fluid is then poured down the thistle-funnel from time to time, the additions being made at regular intervals during the entire procedure. If only a small amount of arsenic be present, the flow of gas must be very gentle, so that if ignited at the concentrated end of the tube little more than a luminous point would be seen. Should the evolution of hydrogen begin to fail, a little more dilute acid must be poured down the thistle-funnel, on no account should strong acid be used, and if the flask becomes at all warm, it should be placed in a vessel containing cold water. The Bunsen flame is kept steadily playing on the tube for at least an hour. The presence of arsenic reveals itself by the gradual formation of a deposit, an inch or more away from the flame, in the direction of the free end of the tube. In its most typical form the deposit appears as a lustrous, nut-brown coating, beginning where the tube narrows and extending half an inch or more along the narrowed part. If the deposit is heavy, it will probably be almost black at the densest part, the margins being brown. On volatilising the deposit, after the tube is detached from the flask, the arsenic combines with oxygen and is redeposited at a little distance from the source of heat, in the form of crystals of arsenious oxide, which may be tested as previously described. On the other hand, if the gas contains but a trace of arsenic, and especially if it is not thoroughly dry, the deposit may take the form of a greyish-white cloud which, under the microscope, is seen to consist of crystals of arsenious oxide.

<sup>1</sup> *Quarterly Journ. of the Chem. Soc.*, 1861

<sup>2</sup> *Mem. Lit. and Phil. Soc. Manch.*, 1904



On heating this grey deposit in a current of sulphuretted hydrogen a yellow sublimate of arsenious sulphide is produced, this distinguishes it from a whitish cloud often seen in the exit-tube, which probably consists of sulphur

When the liquid to be tested contains a larger amount of arsenic, the gas may be ignited as it issues from the exit-tube, in this case it is best to have the end of the narrow part of the tube bent upwards. If much arsenic be present, the flame will be of a whitish-lilac colour, and on allowing it to play on a piece of cold porcelain, such as the lid of a crucible, a brown or black deposit of arsenic is obtained. A drop of a solution of bleaching powder allowed to fall on such a deposit immediately dissolves the portion on which it falls, leaving visible a white circle of porcelain. Ammonium sulphide applied in the same way merely breaks up and partially detaches the film from the porcelain, it dissolves but little. If a deposit is treated with a few drops of nitric acid and heated, it is converted into arsenic acid, after driving off the free nitric acid and allowing the porcelain to cool, a drop of silver nitrate solution, added, produces a brick-red colour of silver arsenate.

Marsh's test may be used with some forms of organic matter which contain arsenic, but with many organic liquids it is impracticable to do so on account of the uncontrollable frothing which takes place, and, further, some of the arsenic may not be set free. Marsh's test is not applicable to liquids containing nitrates, nitrites, chlorides, free chlorine, and sulphur-compounds such as sulphurous acid and sulphuretted hydrogen.

A **biological test** for arsenic has been proposed, founded on the property possessed by certain moulds of developing volatile products with arsenic, which reveal their presence by a garlic-like odour. Gossio,<sup>1</sup> the first to propose this test, found that the most active mould in this respect is *Penicillium brevicaulis*. A convenient plan is to put the substance to be tested, finely divided if solid, into a flask along with some small pieces of bread, or of biscuit, and to sterilise the contents of the flask for half an hour at 120° C. When cold, the bread is inoculated with a culture of *P. brevicaulis*, and the flask is kept at a temperature of about 37° C. If arsenic is present, a garlic-like odour will be developed in twenty-four, or more, hours, it is doubtful whether this is due to the formation of arsenuretted hydrogen or to an organic combination of arsenic. The test is so delicate that it will detect the presence of one-thousandth of a milligramme of arsenious oxide. It cannot be used quantitatively, however, nor is it suited for medico-legal purposes.

Arsenic may be separated from organic matter by the method of Fresenius, described on p. 350. If Schneider's method, founded on the volatility of arsenious chloride, is preferred, the substance to be examined should be finely divided and thoroughly dried in a hot-water oven. When sufficiently dry, it should be powdered in a mortar and then placed in a flask connected with a condenser, the lower end of which dips into a receiver containing a little water, both condenser and receiver should be kept cold with a stream of water. The powdered material is then well covered over with pure strong hydrochloric acid, and is allowed to digest for an hour, after which, by means of a sand-bath, moderate heat is applied to the flask until about three fourths of the hydrochloric acid has passed over. The source of heat is now temporarily withdrawn, and more hydrochloric acid is added to the flask, a fresh receiver being substituted for the one already used, and distillation is resumed. The whole of the arsenic usually comes over by the time the second distillation is complete, if not, the process must be repeated. By this method arsenic is separated from all other metallic poisons with the exception of antimony, and, possibly, tin. Much care is required to avoid accident from "bumping" in the flask, which is specially liable to occur with certain organic substances. The temperature of the sand bath should be carefully regulated so that the distillate may be as free as possible from organic matter.

<sup>1</sup> *Giorn. dell'Acad. di Med., Torino*, 1892.

Another method is to destroy the organic matter with nitric and sulphuric acids. Gautier's<sup>1</sup> improved process affords the best means of carrying this out—100 grms of the fresh tissue, finely divided, are placed in a large evaporating basin and from 20 c.c. to 40 c.c. of nitric acid and 1 c.c. of sulphuric acid are added. With moderate heat the mass liquefies and then becomes viscid, when the heat is withdrawn and 4 or 5 c.c. of sulphuric acid are admixed and the heat is reapplied. After a time the heat is again withdrawn, and about 10 c.c. of nitric acid are cautiously added, drop by drop. Nitrous fumes are given off, and when the reaction is over the mass is strongly heated until the sulphuric acid emits a dense vapour and there remains a dark liquid which is insusceptible of further carbonisation. After cooling, a little more sulphuric acid is added, and the mixture is well stirred and then poured into 600 c.c. to 700 c.c. of distilled water. A humid substance falls to the bottom and the supernatant liquid, dark brown in colour, is filtered off.

Of these methods, that of Fresenius is the best for general use, especially with large amounts of organic matter, if properly conducted there is no risk of loss of arsenic, as has been alleged. Distillation with hydrochloric acid gives excellent results in certain cases, but it is of limited application, some viscera, such as the liver, are troublesome to deal with, and the resulting distillate is usually considerably contaminated with organic matter. Gautier's method is only suitable for small quantities of organic matter, it is not without risk to the operator, nor are the results obtained as perfect as is claimed.

**Quantitative Estimation.**—The filtrate obtained by the methods of Fresenius or of Gautier, or the distillate obtained by Schneider's process, according to the procedure that has been adopted, is placed in a conical precipitation flask, the distillate requires diluting with distilled water. The contents of the flask are raised to about 45° C., and a stream of washed sulphuretted hydrogen, free from arsenic, is passed through the liquid for eight or ten hours, the temperature being maintained all the time. When the liquid is completely saturated with  $H_2S$ , the flask is lightly covered and is set aside until the whole of the precipitate has subsided, this takes one or two days, and its completion may be recognised by the supernatant liquid being perfectly clear and bright. The precipitate, consisting of arsenious sulphide largely contaminated with organic matter and free sulphur, is filtered off, washed with sulphuretted hydrogen-water until free from chlorides, and is then digested in dilute ammonia water (1:15). The solution obtained is filtered and the filtrate is slowly evaporated to dryness. The residue is drenched with nitric acid and dried at a moderate heat, and is then treated with a little sulphuric acid, after which it is heated strongly until fumes of sulphuric acid are given off. By this treatment the precipitated arsenic is converted into a soluble form and the sulphur and the organic matter are oxidised.

If there is sufficient arsenic to be weighed as sulphide, the product last obtained is dissolved in water acidulated with hydrochloric acid and is reprecipitated with sulphuretted hydrogen. The precipitate is collected on a tared filter, is washed successively with absolute alcohol, ether, and carbon bisulphide, so as to remove any free sulphur, is then carefully dried at 100° C. and weighed. 100 parts correspond to 60.98 of the metalloid. After weighing, the sulphide may be reduced to the metallic state by mixing it with potassium cyanide and sodium carbonate, the mixture being placed in a piece of hard glass-tubing narrowed down for a couple of inches at the end. Dry  $CO_2$  is passed through the tube from the thick end, and gentle heat applied until the tube and its contents are free from moisture, the heat is then increased so as to reduce the sulphide to metallic arsenic, this is deposited in the narrow part of the tube, which should be kept cool. The part of the tube containing the deposit may be sealed off from the rest and preserved as proof of the presence of the poison.

If the amount of arsenic is too small to be estimated as sulphide, the product which is left after treatment with sulphuric acid and heat is dissolved in a little water and is estimated by Marsh's method. Quantitative estimation by Marsh's process may be accomplished in two ways—(1) By cutting off the portion of the tube in which the arsenic is deposited and weighing it, the deposit is then dissolved in nitric acid and the tube is dried and reweighed, the difference represents the weight of the arsenic. 75.78 parts of arsenic equal 100 parts of  $As_2O_3$ . (2) By comparing the deposit with the deposits in a series of tubes, of the same diameter, which have been prepared from various known amounts of  $As_2O_3$ . An excellent description of this method is given by Sanger in the *Proceedings of the American Academy of Arts and Sciences*, 1891, and by W. Thompson in the *Manchester Memoirs*, 1903.

### ANTIMONY.

The preparations of antimony, which are met with in toxicological inquiries, are antimony and potassium tartrate, and antimony chloride, chiefly the former.

<sup>1</sup> *Comptes Rendus*, 1899.

**Antimony and Potassium Tartrate** ( $\text{KSbC}_4\text{H}_4\text{O}_7, \text{H}_2\text{O}$ ), or **tartar emetic**, is a well-known medicinal preparation, which contains about 35 per cent of metallic antimony. It is very soluble in water.

**Symptoms of Acute Poisoning.**—When a poisonous dose is taken into the stomach an astringent metallic taste is usually experienced almost immediately, followed in a few minutes by violent pain from the mouth down to the stomach, the pain is hot and burning, and is accompanied by a sensation of constriction in the throat. Immediately after, profuse vomiting comes on, and a little later diarrhoea, blood may be present in the vomit, but more usually is absent. The depressing effects of the poison quickly show themselves in the form of a small, frequent pulse, diminished arterial tension, cold clammy surface, shivering, and profound collapse, in this stage the surface may be cyanosed and the patient unconscious. Clonic spasms may precede death. Respiration is slow and laboured. The urine may be almost entirely suppressed.

Anomalous symptoms not unfrequently occur. Vomiting may be delayed as long as an hour after the poison is swallowed, and then it may either be slight or very violent. In some instances the symptoms resemble those produced by a narcotic. Dobie<sup>1</sup> records a case in which one drachm of tartar emetic was followed by a comatose condition, the patient dying on the sixth day. Respiration is not invariably affected, Carpenter<sup>2</sup> records a case in which, after 170 grains dissolved in water were swallowed, the respirations remained unaffected, recovery taking place.

**Fatal Dose.**—The smallest recorded is one grain and a half of tartar emetic, following a similar dose taken twenty-four hours previously. No effect was produced by the first dose, but the second caused violent vomiting and purging, death taking place in about thirty-six hours, the patient was a healthy woman of twenty-five years of age.<sup>3</sup> This is an exceptional case. Five to ten grains probably represent a minimum fatal dose for a healthy adult. Children have succumbed to less. On the other hand, recovery in adults has followed 170 grains (see above) and in one case even 200 grains. Death may occur in from a few hours to several days.

When large doses are quickly rejected, the local effects of the poison are often speedily recovered from, the danger to a great extent appears to lie in its depressing effects, including those due to violent vomiting and purging.

**Sub-acute or Chronic Poisoning.**—When death results from homicidal poisoning with tartar emetic it is usually due to repeated doses, which progressively depress the systematic powers, prevent the retention of food, cause persistent vomiting and purging, and thus eventually lead to a fatal issue.

In the case of *Reg v Pritchard* (High Court of Justiciary, 1865), the prisoner was accused of having poisoned his mother in law and his wife by the administration of tartar emetic. His wife was in her usual health up to the end of October, 1864, when she began to suffer from frequent attacks of vomiting, on leaving home she gradually regained her usual health, but after her return she again began vomiting and was attacked with severe cramps. The vomiting occurred within an hour or two after meals, which were always sent to her by her husband, not only food but also beverages, such as camomile tea, egg flip, and port-wine were rejected. Death took place on March 18, 1865. During the illness of the deceased cramps constituted a prominent symptom, the wrists were turned in and the thumbs powerfully flexed. Retching, vomiting, and diarrhoea were persistent, the tongue was foul, and there was constant thirst with profound depression. A considerable amount of antimony was found in the viscera after death, especially in the liver and in the contents of the intestines, in which it was present in a soluble form, it is probable that the last dose of the

<sup>1</sup> *The Lancet*, 1887.

<sup>2</sup> *New York Med Rec*, 1883

<sup>3</sup> *Bulletin de Therapeutique*, vol. II.

poison was administered a short time before death. The prisoner was condemned, and, before his execution, confessed that he had poisoned the deceased.

The case of *Rez v Kłosowski*, described on p. 334, furnishes other instances of homicidal poisoning by the frequent administration of small doses of tartar emetic.

Tartar emetic has been administered in dangerous doses to drunkards during a debauch, not with the object of injuring them, but to produce such vomiting and nausea as to render them incapable of further excess for the time being. In a case of this kind seen by the author, the patient vomited and was purged excessively, the tongue was foul and the surface was cold, but the pulse and respiration were not much affected. Cramps of the muscles of the limbs were severe and continuous, they lasted more than forty-eight hours, and persisted in the hands after the other parts originally attacked were free. Antimony was found in the urine. The patient recovered from what in all probability was a single dose, with a celerity that afforded a striking contrast with the severity of the symptoms.

**Antimony Chloride** ( $\text{SbCl}_3$ ), or butter of antimony, used in an impure state for certain trade purposes, has exceptionally been administered as a poison. In addition to the toxic effects of antimony, this salt has a powerfully corrosive action on the tissues with which it comes in contact and produces corresponding symptoms and post-mortem signs.

**Treatment of Acute Antimonial Poisoning.**—The poison usually promotes its own evacuation, if it has not done so, the stomach must be emptied either with the stomach-tube or an emetic, or probably tickling the throat will be sufficient to excite copious vomiting, when antimony chloride has been taken the stomach-tube should be used with great caution, if used at all. Then tannin, or some substance containing it, should be given so as to form an insoluble combination with any of the poison that remains in the stomach. After the poison is evacuated, useless vomiting should be checked by ice and opium. External warmth should be applied, and stimulants given if needed. It is characteristic of antimony poisoning that, after a large dose, either a speedily fatal issue from exhaustion may be expected, or an almost equally speedy recovery, in this respect it differs from arsenic. Antimony is eliminated by the kidneys and the bowels.

**Post-mortem Appearances.**—After acute poisoning by tartar emetic, the mucous membrane of the **stomach** is usually strongly injected and swollen, and in parts may show indications of loss of superficial portions, it is covered with a slimy mucus, and is frequently ecchymosed. A similar appearance, less intense, will be found in the duodenum. In some instances the mucous membrane of the stomach has been found ulcerated, and detached from the muscular coat, the mucous membrane of the œsophagus being also ulcerated, in others there is no appearance of ulceration of the stomach, the mucous membrane being also free from signs of inflammation. In the case of Bravo, who was poisoned with tartar emetic in 1876, both stomach and duodenum were pale and yellowish on their inner surfaces, there were ulcers in the cæcum, and the large intestines were blood-stained. The mucous membrane may be stained orange yellow, due to the formation of antimony sulphide. The **liver** and **kidneys** may show fatty changes, usually only when death has been caused by several isolated doses. The post-mortem appearances of poisoning by antimony are neither so characteristic nor so constant as in arsenical poisoning (See results of post-mortem examinations in the cases of poisoning by tartar emetic, by Stevenson <sup>1</sup>).

<sup>1</sup> *Brit. Med. Journ.*, 1903.

Antimony chloride produces post-mortem appearances like those of a corrosive, such as hydrochloric acid, exceptionally, corrosive effects may be absent. Cooke<sup>1</sup> records a case of a woman, aged forty, who immediately after a meal swallowed the contents of a four-ounce bottle of butter of antimony, vomiting without blood occurred, and profound collapse, death took place in less than two hours. On post-mortem examination no corrosion of tongue, mouth, fauces, or œsophagus was found, the mucous membrane of the stomach was intensely congested, almost black.

**Chemical Analysis—Tests.**—Organic fluids, or solids, when pulvified and mixed with water to a fluid consistency, may be tentatively examined by **Reinsch's method**, as described in the preceding section. The amount of antimony present, and the length of time the solution is boiled with the copper-foil, and, to some extent, the acidity of the solution, determine the appearance presented by the deposit, if it is but slight, the foil merely acquires a purple tint, if thicker, it resembles tarnished sheet zinc, and if very copious, it is covered with an amorphous black coat. When the foil is heated in a reduction-tube, the deposit is volatilised, and condenses on a cooler part of the tube as a white amorphous cloud of antimonious oxide, which, when only minute traces of antimony are present, is scarcely visible. Under the microscope no trace of crystalline formation can be seen, sometimes there is an appearance that at first sight might be taken for a crystalline deposit, but careful examination corrects the impression.

After shaking out the copper-foil the deposits of antimonious oxide may be dissolved with gentle heat in a couple of drops of a solution of tartaric acid, and tested with  $H_2S$ , which gives the orange-coloured sulphide. If preferred, the deposit of antimony on the copper-foil may be dissolved off in a weak solution of potassium hydroxide, to which a little potassium permanganate has been added, heat is applied until the deposit is dissolved, and then the fluid is separated by filtration from the precipitate of manganese that forms, is acidulated with  $HCl$ , and treated with  $H_2S$ .



Fig 25.—Deposit of antimony on exit tube of Marsh's apparatus  
The arrow indicates the position of the Bunsen flame

The suspected fluid may be further tested by **Marsh's process**, as described in the last section. Antimoniuretted hydrogen is without odour and burns with a greenish-white flame.

**Examination of the Deposits.**—In thin films the deposit on **porcelain** obtained from the flame is of a neutral tint, without trace of brown, where the deposit is heavy, it is amorphous and black—like a smoked surface. The deposit is insoluble in a solution of bleaching-powder, but is freely soluble in ammonium sulphide, leaving, when dry, an orange-coloured residue of antimonious sulphide. When a deposit is dissolved in  $HNO_3$  with heat, evaporated, and treated with a solution of silver nitrate, no colour change is produced, the deposit of arsenic similarly treated yields a brick-red colour.

In the **exit-tube** of the Marsh apparatus the deposit of antimony in the first instance appears immediately over the flame, it then separates into two

<sup>1</sup> *The Lancet*, 1883.

portions, the larger one being towards the free end of the tube, that to the flask-side of the flame being lighter, and sometimes scarcely visible. The two deposits are most widely separated at the under part of the tube where it is hottest, they curve over towards each other at the upper and cooler part. When the deposit has taken up its final position, it is much nearer the flame than is the case with arsenic, all this is due to antimony having a higher volatilising point than arsenic. The deposit of antimony has a bright metallic lustre, like that of mercury, at the part where it fades away furthest from the flame—it is smoky, but without trace of brown, as is the case with arsenic. A very slight deposit may be devoid of metallic lustre, being smoky, or greyish if moisture is present. On applying heat to the tube, the film of antimony is volatilised with difficulty, it comes down again, if oxygen is present, as a white amorphous deposit of antimonious oxide.

Antimony may be obtained from **organic admixture** by acidulating the organic fluid with HCl, and placing it in a platinum capsule, in which a small piece of pure tin or zinc is laid, where the metals touch each other a black deposit of metallic antimony is formed in a few minutes, or in an hour or two, according to the amount present. After removal of the fluid and of the tin or zinc, the deposit should be washed and then treated with strong  $\text{HNO}_3$ , with the aid of heat, and the free acid driven off, the residue may be dissolved in strong HCl. If the solution thus obtained is largely diluted with water, the oxychloride is precipitated as a white insoluble salt, which may be dissolved in a solution of tartaric acid, and precipitated with  $\text{H}_2\text{S}$  as antimonious sulphide, recognised by its orange colour. In some instances antimony may be directly precipitated from organic admixture by passing  $\text{H}_2\text{S}$  through it to saturation, after adding a little tartaric acid, and boiling it to ensure the metal being in solution.

When dealing with the tissues it will be necessary to break them up by the moist process, taking the precaution to adapt a condenser to the flask in which the chlorine is evolved, in order to prevent possible loss from volatilisation of antimonious chloride, there is not much risk, however, as antimonious chloride is not nearly so volatile as arsenious chloride.

**Quantitative Estimation**—Through a given proportion of the filtrate obtained after breaking up the organic matter, sulphuretted hydrogen is passed to complete saturation, so as to precipitate the antimony as sulphide. As antimony sulphide cannot be thoroughly dried at  $100^\circ\text{C}$  it is necessary to conduct the drying in an atmosphere of  $\text{CO}_2$ , otherwise at the requisite temperature it would lose sulphur, probably there will also be some free sulphur to get rid of that has fallen from the  $\text{H}_2\text{S}$ . The precipitate, after drying in the ordinary way at  $100^\circ\text{C}$ , should be powdered in an agate mortar, and put into a porcelain boat, which is placed in a hard glass tube through which dried  $\text{CO}_2$  is passing. Heat is applied until all moisture and free sulphur are expelled, the residue is pure  $\text{Sb}_2\text{S}_3$ , 100 parts equal 71.77 of antimony. If free from uncombined sulphur, the sulphide dissolves in strong HCl without leaving any residue.

## MERCURY.

Metallic mercury in bulk has only very exceptionally produced symptoms of poisoning. In a finely divided state, as in blue pill or blue ointment, or in a state of vapour, the toxic effects of the metal are readily produced. The principal poisonous salt is mercuric chloride, much less toxic, and much less frequently encountered in forensic investigations are mercurous chloride, mercuric oxide or red precipitate, and mercurammonium chloride or white precipitate. Mercuric nitrate resembles mercuric chloride in its effects. When pure, mercuric sulphide, or cinnabar, excepting in the form of vapour, is inert. The combinations of mercury with the organic radicles, methyl and ethyl, are virulent poisons.

### Acute Mercurial Poisoning.

**Mercuric Chloride** ( $\text{HgCl}_2$ ), or corrosive sublimate, dissolves in 16 parts of cold water and in three parts of boiling water. It readily combines with albumen, a property upon which its corrosive action depends.

**Symptoms.**—Immediately after a poisonous dose of mercuric chloride in solution is swallowed, an acrid metallic taste is perceived, which is accompanied by a sensation of constriction in the throat. A hot, burning sensation quickly develops, which spreads from the mouth, along the œsophagus down to the stomach. Vomiting of white slimy masses, which are frequently mixed with blood, rapidly follows. The pain radiates over the abdomen, there it assumes a colicky character, and is succeeded by copious diarrhœa accompanied by severe tenesmus, the stools are watery, often blood-stained, and they, as well as the vomited matters, contain shreds of mucous membrane, blood is more constantly present in the vomited matters and in the motions than with poisoning by either arsenic or antimony. The mucous membrane of the mouth and pharynx is white and swollen, that of the larynx being also frequently swollen, making the voice hoarse and the breathing difficult and noisy. The urine is often completely suppressed for twenty-four hours or more, any that may be passed will probably contain albumen, and it may be tinged with blood, in consequence of the diminished metabolism of proteids the amount of urea present is reduced 30 to 40 per cent. Symptoms of profound collapse are present, the surface is cold, moist, and cyanosed, the pulse being small and irregular. There may be severe hiccough, or convulsions. If the victim survives the early symptoms, salivation is likely to appear twenty-four or more hours after the inception of the poison, this is not always the case, even when a large dose has been taken and recovery takes place. Other symptoms of stomatitis may be present. Richiardi<sup>1</sup> records the case of a woman, aged twenty-five, who swallowed about 45 grains of mercuric chloride in 10 per cent solution, in spite of speedy vomiting and the administration of white of egg, diarrhœa and, on the fourth day, gangrene of the mouth and vagina occurred with hæmorrhage from the bowels on the fifth day, followed by collapse and death. On section the whole of the mucous membrane of the large intestine was found to be gangrenous. In a case, recorded by Coates,<sup>2</sup> a man aged forty-two drank a tea-cupful of a solution of mercuric chloride which contained about forty grains, supposing it to be tea. Tightness of the throat, which was red and inflamed, intense abdominal pain, incessant vomiting, followed in three hours by diarrhœa, colic, sweating, and prostration occurred. The teeth became loose, there was salivation, and the gums bled freely. The patient had a constant dull, lumbar pain, and from the first to the eighth day there was total suppression of urine, subsequently, the kidneys again began to act and urine was passed. The teeth became firmer, but the vomiting and diarrhœa continued, and on the eighth day bright-coloured blood was vomited, this continued for four days, when the patient died. At the necropsy, the mucous membrane of the stomach was almost gangrenous, and there was a distinct perforation about three and a half inches from the cardiac end of the viscus. There were patches of incipient gangrene in the intestines. Exceptionally, the onset of the symptoms may be delayed, sometimes inordinately so, as in the following case, reported by Spillmann and Blum.<sup>3</sup> A woman aged thirty drank a solution containing 46 grains of mercuric chloride. For forty-eight hours she presented no symptoms except almost entire loss of vision. Diarrhœa then abruptly commenced, followed by blood in the motions. Salivation and facial œdema occurred, followed, after a passive period of thirty-six hours, by convulsions, in which the patient died twelve days after taking the poison. Post-mortem, intense inflammation of the large intestine with gangrene of the rectum were

<sup>1</sup> *L'Union Med.*, 1896.<sup>2</sup> *La Méd. Moderne*, 1904.<sup>3</sup> *The Lancet*, 1899.

found The stomach was œdematous The kidneys were large and white, but there was no albuminuria (<sup>1</sup>) The amaurosis was probably due to œdema of the retina, and death to uræmia

The external use of corrosive sublimate has produced fatal poisoning from applications of a strong solution to ulcerated surfaces, absorption may also take place through the unbroken skin The extensive use of corrosive sublimate as an antiseptic has occasionally been the cause of accidents Legrand <sup>1</sup> records the case of a woman who received two uterine injections of a solution of corrosive sublimate, 1 in 2,000, and in consequence died in three days Hall <sup>2</sup> describes a similar case in which death occurred on the tenth day In a case related by Huber <sup>3</sup> a woman accidentally had 150 c c of a 0.5 per cent solution of corrosive sublimate, diluted with an equal volume of water, administered as an enema, violent vomiting and purging set in, with subsequent collapse, which ended in death on the fifth day Schildecher <sup>4</sup> reports three fatal cases following the introduction of bichloride tablets into the vagina for the purpose of preventing conception In each case attempts were made to remove the tablets owing to the burning pain produced, and in one a physician arrived within twenty minutes and immediately gave a vaginal injection of hot water followed by hot milk These cases show how rapidly a lethal dose can be absorbed

**Fatal Dose.**—Three grains of mercuric chloride taken by the mouth have caused death in a child From 3 to 5 grains would probably prove fatal to an adult Recovery has followed 90 grains in one case, in which salivation occurred, and 100 grains in another, in which there was no salivation Joulia <sup>5</sup> records the remarkable recovery of a woman who swallowed 370 grains of mercuric chloride with an equal quantity of tartaric acid Incessant vomiting occurred, the tongue, mouth, and pharynx were swollen and dark red in colour, the pulse was 140 The stomach was immediately washed out The following day anuria and diarrhœa occurred the third day ten ounces of blood-stained, albuminous urine containing casts were passed Recovery took place Death usually occurs within three or four days, it may take place in a few hours, or it may be delayed for seven or eight days

**Red Precipitate (HgO),** when taken in large doses, produces the usual symptoms of irritant poisoning Ord <sup>6</sup> relates a case in which one teaspoonful was taken, vomiting, diarrhœa (without blood) and tenderness of the abdomen were present, recovery took place without salivation In another case recovery followed two drachms Mitchell <sup>7</sup> records the case of a man, aged forty-seven, who swallowed an unknown amount of red precipitate which caused pain in the abdomen, vomiting, and diarrhœa, followed by death The mucous membrane of the stomach was found softened and particles of the poison were seen adhering to its surface

**White Precipitate (NH<sub>2</sub>HgCl)**—About 35 grains, sold in mistake as one of the ingredients of a seidlitz powder, caused the death of an adult Forty grains taken by a man, aged fifty-two, caused death in five hours Twenty grains produced violent vomiting and purging with blood in the stools, in a woman of forty-eight, followed by salivation and stomatitis, recovery took place In other cases it has been taken in much larger doses—as much as two drachms—without fatal result

<sup>1</sup> *Ann. de Gynécologie*, 1890

<sup>2</sup> *The Lancet*, 1912

<sup>3</sup> *Zeitschr. f. klin. Med.*, Bd 14

<sup>4</sup> *Amer. Journ. of Obstetrics*, 1911.

<sup>5</sup> *Gaz. Méd. du Centre*, 1899

<sup>6</sup> *The Lancet*, 1888.

<sup>7</sup> *Boston Med. and Surg. Journ.*, 1897



**Mercuric Nitrate** [ $\text{Hg}(\text{NO}_3)_2 \cdot \text{H}_2\text{O}$ ] on several occasions, when administered as a poison, has been in admixture with free nitric acid, in which condition it is used for veterinary purposes, it is a powerful corrosive. Half an ounce of the solution in nitric acid caused the death of an adult in twenty-five minutes. Hall<sup>1</sup> describes a case in which a teaspoonful caused the death of a middle-aged woman on the eighth day. The symptoms were those of mercurial intoxication rather than nitric acid poisoning. Death has also resulted from its external use as an escharotic.

**Potassio-Mercuric Iodide** acts much in the same way as the chloride. Davies<sup>2</sup> records the case of a young man who swallowed twenty-four "soloids," each containing one grain of this double salt, dissolved in water. Pain in the throat and abdomen quickly followed, with vomiting and moderate purging, the vomit contained some blood. Under treatment, recovery rapidly took place. Salivation was not observed.

Metallic Mercury in a finely divided state is absorbed as such, but is probably partially oxidised, and combined with albumen before being taken up by the blood. Mercuric salts, as before stated, at once combine with albumen, and probably exist in this combination in the systematic fluids, being held in solution by excess of albumen. Mercurous salts are taken up with difficulty on account of their insolubility, a large proportion being rejected either by vomiting or by the bowels, the relative insolubility of mercurous salts does not deprive them of toxic properties, however, as is shown by many instances of fatal poisoning with them. Runeberg<sup>3</sup> relates the case of a woman who received three subcutaneous injections of calomel— $1\frac{1}{2}$  grains in each—within one month. Ulcerative stomatitis, with profuse salivation and diarrhoea, set in, followed by collapse and death, which took place in a few days.

Mercury is **eliminated** by the urine, faeces, saliva, and skin. When abundantly present in the body it may be found in the serum of blisters, in the milk, and in any other normal or abnormal secretion.

**Treatment.**—In acute poisoning, if emesis has not already occurred, the stomach should be emptied by an emetic followed by plentiful administration of raw white of egg, the albuminate thus formed, although insoluble in water, is soluble in excess of albumen, therefore it should be removed as quickly as possible by producing further vomiting. Magnesium carbonate is useful to reduce the mercuric salt to a less active form. Afterwards demulcents and opium will be required.

**Post-mortem Appearances of Acute Mercurial Poisoning.**—Taking the appearances of mercuric chloride as a type, the lips and the mucous membrane of the mouth, including that of the tongue, are usually swollen, softened, and of an ash-grey or white colour, this appearance may persist along the oesophagus, the affected membrane being sometimes corrugated, and sometimes eroded. The mucous lining of the **stomach** is swollen and softened, it has been found deeply injected—of a bright scarlet colour—with ecchymoses, in other cases the indications of inflammation are not nearly so obvious, eschars have been found in the vicinity of the pylorus. The small intestines are usually much less affected than the caecum, colon, and rectum, which are generally deeply injected, the lining membrane being probably ulcerated in parts, with indications of hæmorrhage, if death is very rapid, the intestines may present no abnormal appearance. Evidence of interstitial **nephritis** will probably

<sup>1</sup> *The Lancet*, 1912.

<sup>2</sup> *Brit. Med. Journ.*, 1907.

<sup>3</sup> *Arch. / Dermatologie u. Syph.*, 1889.

show itself, unless the case was so very acute as not to allow time for its production. Deposits of lime salts have been found in the tubules of the cortex of the kidneys. A case was investigated by Kaufmann,<sup>1</sup> in which a woman of twenty years died in nineteen days after swallowing a solution containing from 8 to 12 grms (124 to 186 grains) of mercuric chloride. On the fourth day anuria occurred, which lasted two days, on the three following days the urine was plentiful. On section the kidneys were found to be deeply injected and to contain a number of calcareous deposits in the cortices. Microscopical examination showed that the appearances presented by the kidneys did not depend on parenchymatous inflammation, but on a non-inflammatory necrosis of the epithelium—coagulation or anæmic-necrosis, the calcareous deposit existed in the epithelial cells and not in the lumen of the tubules. The epithelial layer of the mucous membrane of the stomach also contained numerous minute calcareous deposits. A number of thromboses due to alteration of the blood, which is regarded by Kaufmann as the essential nature of corrosive sublimate poisoning, were found in the capillaries of the lungs and elsewhere, but whether the blood stasis in the capillaries is caused by changes in the red corpuscles themselves, or by liberation of fibrin-ferment, is not certain, possibly by both.

### Chronic Mercurial Poisoning.

When mercury is taken into the system in repeated small doses, the effects produced are of a special character, differing in their salient features from those met with in acute poisoning. Chronic mercurial poisoning occurs almost exclusively among workers in the metal or among those who handle substances which contain its salts—looking-glass makers, thermometer and barometer makers, workers in quicksilver mines, and in manufactories in which the preparations of quicksilver are produced, represent the first division, furriers, bronzers, and others the second.

The **order** varies in which the **symptoms** of chronic mercurial poisoning occur. The first indications usually are symptoms of dyspepsia, anorexia, colicky pains, loss of flesh and of strength. Increase in the secretion of saliva is observed, accompanied by fœtor of the breath, tenderness of the gums, and the general symptoms of stomatitis. The sufferer looks anæmic, he is subject to attacks of nausea, vomiting, and diarrhœa. The skin shows erythematous, eczematous, or pustular eruptions. Exceptionally, glycosuria occurs, either in cases of accidental poisoning with mercury, or even after its therapeutic use. Bing<sup>2</sup> records the occurrence of several cases of chronic mercurial poisoning caused by the derangement of a mercury-ventil in a hospital. The symptoms comprised shortness of breath, cyanosis, and nausea, followed by vomiting. Two of the cases ended fatally, section showed hyperæmia of the lungs, in which the smallest bronchioles were affected, and atelectasis. A like condition was produced experimentally in animals, by means of allowing them to breathe the moist air which had passed through mercury. The effects thus produced differ from those encountered in ordinary chronic mercurial poisoning.

Sooner or later the occurrence of special symptoms indicate that the nervous system is implicated, in some instances the nerve-symptoms are the first to show themselves. The earliest and most characteristic is a fine tremor of the muscles of the tongue and face, at first only manifest under excitement, the tremor tends to spread to the arms, and later to the legs. Although at first

<sup>1</sup> Virchow's *Arch.*, 1889

<sup>2</sup> *Arch. f. Hygiene*, 1903.

called up by exertion only, its fineness—resembling paralysis agitans—distinguishes it from that of disseminated sclerosis. Subsequently the tremors are continuous, although still accentuated on voluntary movement, rendering co-ordinated muscular action difficult, during sleep they may be absent, or simply lessened, in accordance with their intensity when the patient is awake. As is the case in all tremors that affect the muscles of articulation, the patient stammers and hesitates when speaking. The tremors may exist without appreciable loss of muscular power, but, as a rule, more or less paralysis occurs. Letulle<sup>1</sup> found this to be the case with a number of workers in the Almaden mercury mines, the diminution in muscular power, as tested by the dynamometer, was proportional to the duration of the mercurial influence. Muscular weakness may occur without tremors, but complete paralysis is invariably preceded by tremors. Sensory disturbances, such as imperfect tactile sensibility, hyperæsthesia, and painful sensations, are, as a rule, localised and not profound. Psychical disturbances are frequent, they take the form of mental irritability and loss of power of concentration, with headache and palpation, the condition called “mercurial erythism” may be present, the patient being subject to hallucinations and to acute attacks of mania. In the majority of cases observed by Letulle, the digestive organs were healthy. The teeth may be blackened, and appear as though corroded by an acid, the condition, however, differs from ordinary caries.

**Chemical Analysis.**—The presence of mercury in organic admixture, in not too small amount, may be demonstrated by Reusch's test. The film of mercury on the copper-foil is very characteristic, presenting the appearance of polished silver, if but a trace be present the foil is barely discoloured. The foil, after being dried, is put into a reduction-tube and heated until the mercury is driven off, when minute globules of the metal are deposited on the cooler part of the tube. Under the microscope by transmitted light they appear like black balls, by reflected light they show a metallic lustre round their margins. When the tube is cool (the copper being shaken out), the vapour given off from a scale of iodine introduced into it soon colours the mercurial deposit yellow, which gradually deepens into scarlet mercuric iodide. The presence of mercury in organic fluids may also be ascertained by immersing in the fluid, after acidulation with hydrochloric acid, a slip of gold-foil in contact with a piece of tin wire, a white stain (metallic mercury) appears on the foil where it is touched by the tin. On account of the volatility of the metal the moist method should be used in order to separate mercury from organic matter. If the amount of mercury is not very small, the liquid obtained after treatment of the organic matter with potassium chlorate and hydrochloric acid may be saturated with  $H_2S$ , and allowed to stand until a black precipitate of mercurous sulphide falls, which after separation must be thoroughly washed free from all trace of chlorides. Any silver, lead, or copper sulphides present may be separated by treatment with nitric acid, in which they are soluble, mercurous sulphide is insoluble in nitric acid. After the precipitate is washed and dried, it may be weighed and the amount of mercury calculated—100 parts equal 86.2 mercury. The sulphide is then treated with nitro-hydrochloric acid, evaporated to dryness, the residue dissolved in water, and the solution tested in various ways for mercury.

**Tests.**—With mercuric salts potassium iodide gives a scarlet precipitate soluble in excess. Potassium hydroxide gives a yellow precipitate. Stannous chloride gives a white precipitate of mercurous chloride which changes to grey—metallic mercury. With soluble mercurous salts potassium hydroxide gives a black precipitate, potassium iodide a green precipitate, stannous chloride a white precipitate changing to grey, potassium chromate a brick-dust coloured precipitate.

A number of methods have been devised for estimating mercury when present in very small amount in organic matter, some modification of the moist method is usually adopted

<sup>1</sup> *Arch de Phys. norm. et pathol.*, 1887.

to destroy the organic matter, and much ingenuity has been expended on the subsequent separation of the metal. Hofmeister's method as adopted by Winternitz<sup>1</sup> dispenses with destruction of the organic matter when urine is the fluid to be examined. The urine is acidulated with 10 per cent of HCl, and is left for two days to deposit uric acid, after filtration it is slowly passed through a system of glass tubes containing rolls of copper gauze, the mercury present is deposited in the metallic state on the gauze, and after washing and drying is driven off by heat, and is deposited in a cool part of the combustion-tube in which the volatilisation is performed, it is then weighed. Bohm<sup>2</sup> modifies this process by destroying the organic matter in the moist way, and after freeing the liquid from chlorine allows it to pass over copper gauze as above described. Ludwig and Zillner,<sup>3</sup> after destroying organic matter with hydrochloric acid and potassium chlorate, precipitate the mercury with zinc dust, from which it is volatilised by heat, the tube containing the deposit is weighed, and again after the mercury is driven off by heat.

These recent methods, which for the most part are modifications of older processes, present certain advantages, and enable approximately exact results to be obtained, but after some experience in their use the author prefers the electrolytic method, as being easier of application and of equal accuracy. The organic fluid, after treatment with potassium chlorate and hydrochloric acid, is submitted to electrolysis, as described in the following section, a small slip of gold foil, as the cathode, being substituted for platinum. After the mercury is deposited, the gold foil is washed, first with water, then with absolute alcohol, and lastly with ether, and is carefully dried and weighed, it is then introduced into a piece of hard glass tubing through which a current of dry air is passed, and sufficient heat is applied to drive off the mercury from the foil on to the tube. The foil is reweighed, and, for control purposes, the tube is weighed with the deposit, and again after it has been driven off by heat.

### LEAD.

The salts of lead usually encountered as toxic agents are the neutral acetate (sugar of lead), the basic acetate (Goulard's lotion), the carbonate (white lead), the tetroxide (red lead), and the chromate (yellow chrome), other salts of lead, as the chloride and nitrate, are poisonous, but are not so accessible to the public. Fine particles of metallic lead are poisonous when repeatedly taken into the system.

The salts of lead act as **mild irritants**, some being more powerful poisons than others, the chromate, for example, although insoluble in water, acts more energetically than the acetate, which is soluble.

Lead poisoning may be **acute** or **chronic**, an intermediate **subacute** form is not unfrequently encountered.

### Acute Lead Poisoning.

**Lead Acetate**  $[\text{Pb}(\text{C}_2\text{H}_3\text{O})_2 \cdot 3\text{H}_2\text{O}]$ , the salt of lead most frequently used, has a sweetish taste, it only produces acute poisoning when taken in large doses.

**Symptoms.**—If an ounce or more is swallowed, a strong astringent metallic taste is at once perceived, followed by a feeling of constriction in the œsophagus, and a hot sensation, which spreads to the stomach. Within half an hour after, vomiting comes on, the vomited matter consisting of white opaque masses, which may be tinged with blood. There is great thirst, and violent colicky pains in the abdomen, which come on in paroxysms, the abdominal muscles are tense, and the patient eases the pain by bending himself forward and compressing the abdomen. The bowels are usually constipated, but, exceptionally, diarrhœa has occurred, the motions are dark, almost black, from the presence of lead sulphide. The urine may be partially suppressed. Great prostration, vertigo, and pains in the head and limbs are experienced, with general numbness, or paræsthesiæ, cramps in the calves, and, occasionally, paralysis of the

<sup>1</sup> *Arch f exp Pathol*, 1889.

<sup>2</sup> *Zeitschr. f phys Chem*, 1891.

<sup>3</sup> *Wiener klin Wochenschr.*, 1889.

limbs Drowsiness is not unfrequent, the tongue is coated, and the breath is offensive, the pulse is small and frequent. In acute lead poisoning, from a single dose the gums rarely exhibit the blue line which is characteristic of chronic lead poisoning. The greater number of cases of acute lead poisoning recover. The **subacute form** occurs after taking repeated small, though not minute, doses of a soluble salt of lead. The patient is troubled with intense thirst and a metallic taste. Colic, with retraction of the abdominal muscles, is a prominent symptom, the bowels are obstinately confined. The urine is lessened in quantity. The blue line round the margin of the gums is usually present, the pulse is weak and slow, the tongue is coated and the breath is offensive. There may be some of the more acute symptoms present, as prostration, numbness, and vertigo. Death rarely occurs, the symptoms passing off in a week or two after the poison ceases to be administered. Gastric disorder and colic are the symptoms which first indicate the occurrence of the subacute form of lead poisoning, which occasionally happens in consequence of active medicinal treatment by means of lead acetate. Usually lead acetate may be administered in medicinal doses for a considerable time without producing poisonous symptoms, as, for example, when given to check the obstinate diarrhoea which accompanies tubercular ulceration of the bowels, on rare occasions poisonous symptoms follow a single dose.

**Fatal Dose.**—The exact amount of lead acetate which will cause death is not known. Recovery has followed an ounce. Lesser<sup>1</sup> records the case of a woman aged forty who swallowed a “knife-pointful” of litharge in order to procure abortion. In an hour and a half she vomited milk-like masses, and on the following day she aborted of a four-months foetus. She became icterish, with abdominal pain, profoundly collapsed, and died on the third day after taking the poison. Slight indications of gastric catarrh were found after death. Lead was found in the viscera. He reports another case in which one ounce and a half of white lead caused death on the third day.

**Treatment of Acute Poisoning.**—The stomach should be emptied either by the tube or an emetic unless there has been free, spontaneous vomiting. Sodium and magnesium sulphates should be given in half-ounce doses, dissolved in half-pints of water, dilute sulphuric acid may be substituted, the lead sulphate thus formed should be got rid of by purgation, as, although it is an insoluble salt, it is not entirely harmless. Demulcent drinks, as barley-water, milk, or white of egg, are beneficial, opium may be necessary for the colic and to restrain useless vomiting.

**Post-mortem Appearances.**—On account of the comparative rarity of fatal cases, the post-mortem signs are not well known. In addition to the usual indications of acute gastro-enteritis, the mucous membrane of the stomach has been found to be covered with a whitish-grey deposit. It may also be thickened and softened, a condition which sometimes extends to the duodenum. Erosions of the gastric and intestinal mucous membrane have been observed, apparently due to prolonged localised action of the lead salt. The other organs yield no reliable indications.

### Chronic Lead Poisoning.

The sources from which lead is derived in cases of chronic poisoning are numerous. They may be divided into those due to **occupation-risks**, and those due to the **accidental presence** of lead in fluids and comestibles, and in

<sup>1</sup> *Vierteiljahrsschr. f. ger. Med.*, 1898

substances which are repeatedly brought in contact with the surface of the body, in the latter case it is probable that the poison is accidentally transferred to the mouth by the fingers, in handling food for example

The following are the notifications of chronic lead poisoning in factories and workshops in the years 1909 and 1919 <sup>1</sup>

Industry	1909		1919	
	Total Cases Notified	Deaths	Total Cases Notified	Deaths
1 Smelting of metals,	66	5	24	5
2 Brass works,	5			
3 Sheet lead and lead piping,	9	2	2	1
4 Plumbing and soldering,	28		10	
5 Printing,	21	1	10	1
6 File cutting,	8			
7 Tinning and enamelling,	21		2	
8 White lead,	32	2	10	
9 Red lead,	10		15	
10 China and earthenware,	58	5	21	8
10a Lath transfers,	1			
11 Glass cutting and Polishing,	4	2		
12 Vitreous enamelling,	3		1	
13 Electric accumulators,	27	2	48	2
14 Paints and colours,	39	2	11	
15 Coach building,	95	6	11	3
16 Shipbuilding,	27	1	8	2
17 Paint used in other industries,	42		9	3
18 Other industries,	57	2	25	1
TOTAL,	553	30	207	26

Among the **accidental** causes are —drinking-water which has been stored in lead cisterns, has passed through lead pipes, or is derived from contaminated sources ( $\frac{1}{100}$  of a grain per gallon has caused poisoning) The risk hitherto encountered of drinking-water becoming contaminated in its passage through lead-pipes is now materially increased by the action of stray currents of electricity emanating from subterranean electric light and power cables Latham <sup>2</sup> relates a case of lead-poisoning which was traced to this cause It was found that when a current of 30 amperes was passing through the offending cable there was a difference in potential of 18 volts between the earth-current and that of the water pipe Danger occurs from food cooked in so-called tinned vessels, the coating in the cheaper kinds sometimes containing lead, or in earthenware vessels lined with lead glaze, or food preserved in "tins," the solder of which contains a percentage of lead Wine bottles that have been cleaned by shaking lead shot within them, confectionery coloured with lead chromate, tea and snuff packed in lead-foil, hair dyes and cosmetics containing lead, and soda-water syphons fitted with pewter or lead valves, have all from time to time given rise to chronic lead poisoning No combination of lead can be taken into the system without the risk of chronic poisoning, even lead sulphate,

<sup>1</sup> Reports of Chief Inspector under Factories and Workshops Acts.

<sup>2</sup> *The Electrician*, 1905

which is regarded as an insoluble salt, was effectively used by Gusserow<sup>1</sup> to produce chronic lead poisoning in animals

Lead or its compounds may enter the system through the respiratory or gastro-intestinal tracts, or through the skin, the last portal being relatively unimportant. Workers in metallic lead do not suffer unless they are frequently in the presence of large quantities of the molten metals, or inhale fine particles of solid lead or of its oxide whilst manipulating old metal, ordinary plumbers, who handle unoxidised metallic lead all day long, comparatively rarely suffer from lead poisoning unless from the use of white or red lead in fitting. Lead, though not usually classed amongst the volatile metals, is capable of volatilisation at a high temperature, and in the form of vapour may be taken into the system through the respiratory tract and also into the stomach. One of the worst cases of chronic lead poisoning seen by the author was that of a man who bought the sheet-lead linings of old tea-chests and melted them down into pig-lead, he did the work in a small room without any contrivance for ventilation, and attended to the whole process himself. File cutters, by constant hammering, abrade and receive into the mouth and nostrils small particles from the thick plates of lead on which they embed the files. In other trades in which lead is used, the salts of the metal are introduced into the system through want of cleanliness, a workman is content to wipe his hands on his apron, or other cloth already contaminated with lead, and then to handle food in the act of taking his meals, he will also hold between his teeth brushes or other articles soiled with lead paint. Goadby has shown by experiments on animals that the inhalation of lead dust is far more dangerous and produces symptoms far earlier than the direct ingestion of a very much larger quantity of the same substance through the mouth.

Idiosyncrasy has much to do with chronic lead poisoning, of half a dozen persons subjected to the same risks, perhaps one only develops symptoms. The use of alcohol increases the tendency to chronic lead poisoning, Oliver<sup>2</sup> regards it as a most potent factor. Gouty subjects easily succumb to the influence of lead, and in its turn the metal tends to develop gout. Oliver states that women succumb to the influence of lead more quickly than men at an earlier age—18 to 23, in the case of men the usual age is from 41 to 48. Lewin<sup>3</sup> states that of one hundred women-workers in the Vienna lead foundries 264 were attacked with saturnism, of the same number of men only 69 were attacked. In the Staffordshire potteries (before the recent regulations were enforced), 40 per cent of women and only 7 per cent of men were affected. Cases have been observed in which symptoms of chronic poisoning appeared yearly after the individual had ceased to be exposed to the action of the metal. In pregnant women abortion frequently results from chronic lead poisoning. According to Lewin, the ecbohic action of lead affects not only women who are suffering from plumbism, but also a healthy woman who is impregnated by a man who, at the time, is suffering from chronic lead poisoning. Abortion from chronic lead poisoning occurs from the third to the sixth month. Pope<sup>4</sup> reports the cases of two women who died from lead poisoning after taking, as an ecbohic, diachylon plaster made into pills, other similar cases are recorded (*vide* p 104).

**Symptoms.**—The early symptoms are usually referred to indigestion, the patient has pains in the stomach or abdomen, which may or may not be evoked, or increased by taking food, the appetite is diminished and the bowels are

<sup>1</sup> Virchow's Arch., vol. xxi

<sup>2</sup> Berliner Klin. Wochenschr., 1904.

<sup>3</sup> Lead Poisoning in its Acute and Chronic Form, 1892

<sup>4</sup> Brit. Med. Journ., 1893.

constipated. A disagreeable, sweetish, astringent taste is experienced in the mouth, and the breath is offensive. The skin acquires an unhealthy colour, at first it is yellowish, subsequently it is anæmic. Round the free margins of the gums a blue line is seen most strongly marked in the upper jaw, where teeth are absent the blue line is also absent. Occasionally, when the teeth are present, no blue line is visible. The line is due to deposition of lead sulphide in the papillæ of the mucous membrane of the gums, small quantities of food containing sulphur cling to the teeth, and in consequence of decomposition slowly give off  $H_2S$ , which combines with the lead present in a state of solution in the gums. Nutrition is interfered with, and the patient consequently emaciates. The pulse is usually slow and of high tension. Parotitis is an occasional late symptom of plumbism.

Patients in this condition often go on for a long time without much change, although they may continue to be under the influence of the poison. The author once examined a number of men who were employed in a large lead works and, in a considerable proportion, found a distinct blue line on the margin of the gums, these men denied that they had ever suffered any symptom of lead poisoning, or that their health had been adversely affected in any way. They had been engaged in the same work for from five to twenty-two years. Usually, however, one or other of the more pronounced symptoms of chronic lead poisoning is developed. They comprise—**colic**, **arthralgia** (pain in the neighbourhood of the joints), **paralysis**, and **encephalopathy** (psychical disturbances).

**Colic** is usually the first to appear, although cases of typical lead paralysis occur in which the patients deny ever having had colic, as a rule, one or more attacks of colic precede the other symptoms. Colic is ordinarily ushered in by recurrent abdominal pains as mentioned in the general symptomatology, but an attack may suddenly occur without any antecedent pain. In most cases the pain radiates round the umbilicus, and is accompanied by tenesmus and retraction of the abdominal muscles, whether the muscles are retracted or not, they are tense and resistant, and the pain is relieved on pressure. Except in rare cases, when there is diarrhœa, the bowels do not act notwithstanding the tenesmus. During the attacks of colic the pulse becomes still slower, and is full and hard. There is no rise of temperature of any significance.

**Arthralgia.**—The pains to which this name is given probably originate in the sensory nerves of the muscles in the neighbourhood of the joints, they may be preceded by shooting or flying pains, or they may come on suddenly without warning. They occur most frequently about the knees, less frequently about the elbows and shoulders. The sensation is that of a boring, tearing pain, which seems to affect the bone itself. The flexors suffer most frequently, as a rule, the small joints are not attacked. The pain may extend to the muscles of the trunk, especially to the lumbar vessels. Contraction and twitchings of the muscles have been observed.

**Paralysis.**—The muscles most frequently attacked are the extensors of the hand and fingers. The order in which they succumb is usually—the extensor communis, the extensores digiti minimi, pollicis longus, carpi ulnaris, radialis, pollicis brevis, and (after a longer interval) the ossis metacarpi pollicis. The supinator longus usually escapes, and shows up in contrast to the wasted muscles. The result of the muscular paralysis is that, when the arms are held out horizontally, with the palms of the hands downwards, the hands drop, and cannot be raised, the condition being known as “**wrist drop**.” Exceptionally, the paralysis begins in the muscles of the upper arm, the deltoid, biceps, and coraco-brachialis being affected, and in this—the upper-arm type—the supinator



ongus is attacked. Both arms usually suffer, though one may be more affected, or further advanced than the other. The interosseous muscles of the hands and those of the ball of the thumb are sometimes specially attacked, producing the "claw-shaped" hand. The legs may be attacked, but usually not until the arms have suffered for some time, the anterior muscles are the first to suffer, the tibialis anticus, like the supinator longus, usually escapes. The muscles of the trunk are rarely affected. Tremor, increased by movement, may occur in the arms.

The characteristics of lead paralysis are - little if any disturbance of the sensory fibres--probably limited to localised patches of anæsthesia if present--with profound trophic changes the affected muscles undergo extreme atrophy, and yield the reaction of degeneration. Lead paralysis is usually regarded as peripheral, but it does not correspond with the distribution of the peripheral nerve supply, in the arm, for example, all the muscles supplied by the musculo-spiral are not affected. Changes have been found both in the nerves and in the ganglion cells of the anterior cornua of the cord.

**Encephalopathy.**—Psychical disturbances usually commence by headache, hizziness, and sleeplessness, there may also be amaurosis, further development may lead to a condition of drowsiness, or to one of excitability, which may be accompanied by hallucinations or wild delirium. Eclampsia is common, especially in women, and involves an unfavourable prognosis, the convulsions may be repeated at intervals for days, the patient remaining unconscious for some time after each attack. Optic neuritis occurs among those who work in lead, especially in girls, cases of this kind are met with in pottery workers. The neuritis may not affect the vision, or it may cause absolute blindness.

If lead is taken into the system for a prolonged period it tends to cause chronic interstitial nephritis, with the presence of albumin in the urine. Albuminuria does not usually occur in the early stage of chronic lead poisoning.

Lead possesses selective properties in relation to the nervous system, which is attacked by it both centrally and peripherally, it is capable of entering into combination with nerve substance, and thus of directly interfering with its function. Blyth<sup>1</sup> chemically examined the brain of a man who had succumbed to the influence of lead, and found an amount of the metal in it equal to 117.1 milligrammes of lead sulphate.

**Treatment of Chronic Lead Poisoning.**—Removal of the patient from the influence of the poison is essential, when dyspeptic symptoms, especially if accompanied by pains in the abdomen, are complained of, the margins of the gums should always be examined. Various remedies, of which the favourite is potassium iodide, have been used to promote elimination of the metal. From the results of some investigations on this subject,<sup>2</sup> the author arrived at the conclusion that potassium iodide does not influence the rate of elimination of lead. In two cases of chronic plumbism, it was given in 15 and 10 grain doses respectively three times a day for a week or ten days, and then stopped for a like period, and again resumed, during the whole period of the experiments the fæces and urine voided in the preceding twenty-four hours were analysed three times a week. The results showed that slow elimination, chiefly through the bowels, was going on the whole time, and that it was not increased when potassium iodide was being taken, in conjunction with which magnesium sulphate was sometimes given. Several other accredited eliminants were also tried with negative results. The only treatment that to a slight degree seemed to increase the amount of lead in the excretions was a combination of hot baths.

<sup>1</sup> Abstract of *Proc. Chem. Soc.*, 1887-88.

<sup>2</sup> *Brit. Med. Journ.*, 1893.

and general massage, with occasional purges. Fresh air, good diet, as much exercise as can be judiciously taken, with hot baths and general massage, are the means chiefly to be relied on to promote cure in cases of chronic lead poisoning.

The **special symptoms** require appropriate treatment — Colic will require opium, arthralgia, hot fomentations, and probably opium, paralysis, local massage and electricity.

The **elimination** of lead from the system takes place chiefly by the bowels, to a much lesser extent by the kidneys. It has been asserted that lead is eliminated by the skin, but the author has never succeeded in obtaining any evidence of this, in the instances of blackening of the skin by baths containing potassium sulphide, which are cited in proof, the discoloration was probably due to some of the metal, derived from external sources, being present in the pores of the skin.

When a medicinal dose of a soluble salt of lead is administered, about half or two-thirds of it passes in an insoluble form directly through the bowels without being absorbed, what remains is gradually eliminated in the fæces and urine, a small percentage being probably retained in the tissues for an indefinite period. In the course of the investigations above mentioned the author administered to a patient 2 grains of lead acetate three times a day for five consecutive days, on the last day of administration the fæces (227 grms) yielded 1762 grm of lead, equal to about 5 grains of lead acetate, on the second day after the administration ceased, 290 grms yielded 1411 grm (about 4 grains) of the acetate, on the fourth day the amount fell to 0053 grm, and on the sixth to 0006 grm, after which there was little more than a trace. The largest amount obtained from the urine in any one day was equal to a little over 1 milligramme of lead, this rapidly fell to less than one-half, and in a few years after there was a mere trace. In each instance the fæces and urine analysed were respectively the excretions of twenty-four hours. It is evident that only a small percentage of lead is absorbed when given medicinally, the instances, therefore, in which symptoms of lead poisoning are produced by medicinal doses demonstrate that it is not essential for much of the poison to be stored up in the tissues, but that what is stored up exists in a very stable form, and probably in intimate combination with them.

The elimination in cases of chronic poisoning was investigated in a similar way, the fæces and urine passed in twenty-four hours were analysed every second or third day, the weight of the fæces and the volume of the urine being noted on each occasion. The results showed that the daily elimination in the fæces was from five to ten times greater than that in the urine, the amount in the fæces varied from 3 milligrammes of metallic lead down to a mere trace, the largest amount obtained from the urine in any one day was 0.9 milligramme.

**Chemical Analysis**—Any lead that comes away with the excretions, or is contained in the tissues after death, exists in combination with organic matter, from which it requires dissociating before it will respond to reagents. If the amount of organic matter with which the lead is combined is small, it may be evaporated to dryness if a fluid, or simply dried if a solid, and then incinerated at as low a temperature as will affect the purpose. The residue is drenched with nitric acid, the acid driven off with a gentle heat, and the nitrate thus formed is dissolved in a little water, filtered, and tested.

**Tests.**—Sulphuretted hydrogen produces a brown or black precipitate in accordance with the amount of lead present. Potassium iodide gives a yellow discoloration or precipitate, the former when the lead is in very small amount,

the precipitate is soluble in boiling water, from which on cooling it crystallises in gold-coloured scales. In making use of this test with minute quantities of lead that have been treated with nitric acid, it is essential that all the free acid should be driven off, otherwise the reagent is decomposed and a yellow colour is produced by the liberated iodine, there are grounds for believing that some of the extraordinary results which have been obtained with potassium iodide, in testing urine for lead, were due to this fallacy. Potassium chromate gives a yellow precipitate. Sulphuric acid gives a white precipitate, hastened, when in very dilute solution, by the addition of alcohol, the precipitate is soluble in ammonium acetate. A lead salt mixed with sodium bicarbonate, and heated on charcoal in the reducing flame of the blowpipe, yields beads of metallic lead incrustated with its yellow oxide.

When minute quantities of lead are present in combination with large amounts of organic matter, the dry process is tedious, difficult to carry out, and uncertain in its results. The plan adopted in the investigations on the elimination of lead above mentioned was as follows.—The urine was evaporated down to the consistence of gruel, and the faeces were mixed with distilled water to a like consistence, they were then each treated with potassium chlorate and hydrochloric acid, as described on p. 350. The filtrate after cooling was placed in a glass cell, the bottom of which consisted of a sheet of vegetable parchment, the cell was immersed to such a depth in an outer cell containing distilled water acidulated with a few drops of sulphuric acid, that the liquids in the inner and outer cells stood at the same level. A piece of platinum foil exposing a surface of about 50 cm. square, constituting the cathode, was submerged in the liquid contained in the inner cell, a similar piece of platinum-foil constituting the anode being immersed in the outer cell, the pieces of foil were so placed as to be opposite each other, separated by the parchment diaphragm. A current of three or four volts was passed from six to eight hours, after which the foil was removed from the inner cell, gently washed and dried. The metallic lead was dissolved off the foil with dilute nitric acid aided with heat, and, after driving off most of the free acid, the solution was decomposed with dilute sulphuric acid, and an equal volume of alcohol was added. It was then set aside for twenty-four hours. The precipitate of lead sulphate was washed with water containing 12 per cent. of alcohol until all the free acid was removed, it was then separated by decantation, ignited and weighed. The amount of lead was calculated from the weight of the sulphate. 100 parts of sulphate contain 68.319 parts of metallic lead.

Whether the moist or the dry process is used, the residue after the primary filtration should be tested for lead, which may exist as sulphate, and remain undissolved. If the original substance contains lead as sulphate, the salt should be dissolved with heat in an aqueous solution of ammonium tartrate, to which a little free ammonia has been added, and then precipitated with sulphuretted hydrogen. 100 parts of lead sulphide contain 68.61 parts of metallic lead. It is better, however, to convert the sulphide into sulphate, by treating it with nitric, and subsequently with sulphuric acids, after which it is ignited, weighed, and the amount of the metal calculated by the factor for lead sulphate.

In place of the electrolytic method, the solution obtained after destruction of the organic matter in the moist way may be precipitated with sulphuretted hydrogen, and the precipitate dealt with as above described, but when the amount of lead is very small, the electrolytic method is much preferable.

## COPPER.

The salts of copper having a distinctive colour and a strongly astringent taste are ill adapted for criminal purposes, still, cases have occurred in which the sulphate and the acetate have been administered with homicidal intent, acute copper poisoning, however, is usually due to accident or to attempted suicide. Metallic copper is slightly if at all poisonous, many cases have happened of accidental swallowing of copper coins, which in some instances have remained within the digestive tract for a considerable time, but, with one exception, no toxic effects are recorded. Rowley<sup>1</sup> relates the case of a girl ten

<sup>1</sup> *Brit. Med. Journ.*, 1894.

years old who swallowed a halfpenny which remained in the digestive tract for six months without giving rise to any symptoms of poisoning Kearney<sup>1</sup> saw an infant nineteen months old who swallowed a copper coin rather larger than a farthing, which was retained nine months and four days and then passed per anum without producing any injurious effects When a soluble salt of copper is swallowed, it is probably transformed into an albuminate, and, if present in small amount, produces but slight local changes, if in larger amount, it not only combines with any free albuminoid substances in the stomach, but it also attacks the mucous membrane and erodes it

### Acute Copper Poisoning.

The salts that are usually answerable for acute poisoning are the **sulphate**  $[\text{CuSO}_4, 5(\text{H}_2\text{O})]$  or blue vitriol, and the **basic acetate**  $[\text{Cu}(\text{C}_2\text{H}_3\text{O}_2)_2, 2\text{CuO}]$  or verdigris

**Symptoms.** When a poisonous dose of either of these salts is taken, the usual effects of an irritant poison are produced within five or ten minutes There is violent vomiting and purging, pain in the stomach and abdomen, a metallic taste, thirst, and the symptoms of collapse The vomited matter at first is green or blue, the lips also and the inside of the mouth may be thus coloured The vomit may be distinguished from bile by the addition of ammonia-water, which strikes a deep blue with the copper salt, the colour of bile remaining unchanged Pain in the head is frequent, and sometimes convulsions occur The urine is diminished in amount, and may contain blood Jaundice has been observed In children the nervous system from the first may be seriously disturbed, producing profound depression, irregular respiration, tonic or clonic spasms of the muscles of the limbs, or complete paralysis, the condition rapidly passing into coma and death Toxic absorption of copper salts may take place through the defective skin A man applied a strong solution of copper sulphate (containing about 80 grains) in milk to his head in order to cure eczema In twenty-four hours severe symptoms of poisoning—gastro-enteritis—occurred, and in the vomit was copper in large quantities (Spaunbauer)<sup>2</sup>

**Fatal Dose** is not exactly known One ounce of the sulphate, and the same quantity of the acetate, have each proved fatal Death may occur in a few hours, it is more usually delayed for several days

**Treatment.**—If vomiting is taking place it may be aided by draughts of warm water, in which the white of egg is beaten up If necessary, the stomach-tube must be used Demulcents, such as barley-water, arrowroot-water, and milk should be given Morphine may be necessary to relieve pain and subdue useless vomiting

**Post-mortem Appearances.**—Indications of the effects of an irritant will be present probably from the mouth down to the stomach and bowels, the mucous membrane will be tumefied and softened, and in the stomach it may be eroded, the whole tract will show signs of inflammation There may be distinctive indications of the poison, especially with the acetate, in the form of green-coloured particles adhering to the gastric, or intestinal mucous membrane, with the sulphate (if the discoloration exists) the appearance will be that of a bluish stain, the staining may be distinguished from that due to bile by the addition of ammonia-water The liver may show fatty changes

<sup>1</sup> *The Lancet*, 1893

<sup>2</sup> *Wien. med. Wochenschr.*, 1904.

### Chronic Copper Poisoning.

According to the popular idea, copper in the smallest doses is a virulent poison, the least contamination of food with the dreaded verdigris is regarded as being pre-eminently dangerous. Formerly, toxicologists shared these views and never questioned the existence of chronic copper poisoning, modern authorities, on the other hand, with few exceptions, state that it is unknown. It is asserted that workers in the metal never suffer from chronic copper poisoning, according to Pecholier and St Pierre (quoted by Gautier)<sup>1</sup> even those engaged in the manufacture of verdigris are not affected by it. Merkel<sup>2</sup> states that in bronze-powder manufactories, where the workmen are coated all over with the powder—which consists chiefly of copper—no case of copper poisoning has ever occurred. Suckling,<sup>3</sup> however, observed in brass workers wrist-drop and other symptoms indicative of peripheral neuritis, a green line was present on the margins of the gums and the corresponding parts of the teeth were stained green. In other cases metallic taste, dyspepsia, attacks of vomiting, and diarrhœa, with colic, have been observed. It is stated that the colic differs from lead colic in not being accompanied by retraction of the abdominal muscles, the tendency to diarrhœa is also a distinguishing feature from chronic lead poisoning. Some observers who have noticed the line on the margin of the gums describe it as being of a reddish-purple, others as indistinguishable from the blue line of lead, others again as in the cases instanced above. Bernatzik<sup>4</sup> states that the line on the gums is not due to copper which has been absorbed, and, therefore, it is not comparable with the blue line of lead, but that it results from the external deposition, at the junction of the teeth and gums, of minute particles of copper which are acted on by the fluids in the mouth and so made to yield a green or blue colour, when the line is reddish-purple it is due to chronic inflammation of the gums. If the green or the blue line is touched with a solution of potassium ferrocyanide it changes to brown.

Filehne<sup>5</sup> maintains that in animals he has produced genuine chronic copper poisoning manifested by changes in the blood, anæmia, fatty degeneration of the liver cells, proliferation of the interstitial tissue of the liver which leads to a condition resembling biliary cirrhosis, together with degeneration of the epithelial cells of the renal tubules. He holds that from the toxicological standpoint there is a complete resemblance between the action of copper and that of the other heavy metals.

For the present, whilst allowing the possible existence of chronic copper poisoning, it may be accepted that the cumulative action of minute doses of copper is infinitely less injurious to human beings than is the case with similar doses of lead.

Copper may find its way into the system along with food, with which it has been accidentally, or purposely, mixed. Accidental admixture occurs from the use of brass cooking utensils, the risk being increased by want of cleanliness. Certain foods, or condiments, are more liable to act on metallic utensils than others, fats easily decompose, and their acids, and also the vegetable acids contained in some fruits, quickly attack copper. Cases of acute irritant poisoning which are reported from time to time as being due to food which has been cooked or stored in copper vessels, are probably almost invariably instances of toxine poisoning from decomposition-products present in the food,

<sup>1</sup> *Le Cuivre et le Plomb*, 1883.

<sup>2</sup> *Münchener med. Wochenschr.*, 1891.

<sup>3</sup> *Brit. Med. Journ.*, 1888.

<sup>4</sup> *Realencyclopædie d. ges. Heilk.*, xi, 1887.

<sup>5</sup> *Deutsche med. Wochenschr.*, 1895.

the amount of copper needful to produce the violent symptoms of poisoning described in such cases is much greater than could possibly be present, if, indeed, any amount of copper could cause symptoms comparable with those of a virulent toxine. In large establishments it is sometimes customary to substitute copper hot-water pipes for those made of lead, as being ultimately more economical, if water conveyed by these pipes is examined, copper in small amount may be detected, the copper boiler and hot-water cylinder, in ordinary household use may be sources of contamination, especially when supplied with some kinds of water.

Copper is purposely added to certain articles of diet in order to improve their appearance, this is frequently done in the case of preserved green peas and some kinds of pickles, as gherkins. The chlorophyll, to which the bright green colour of the vegetables is due, is partially decomposed with corresponding loss of colour during the process of conservation, and a soluble copper salt (usually the sulphate) is added in order to restore the colour and make them attractive to the eye. Tschirch's<sup>1</sup> investigation led him to the conclusion that the green colour, imparted by copper to preserved vegetables, is due to combination of the copper with phyllocyanic acid— a derivative of chlorophyll— forming copper phyllocyanate, when the whole of the phyllocyanic acid in the vegetable is combined with copper any excess of the metal goes to form copper leguminate which, in its pure state, is blue rather than green in colour. The amount of copper varies in different specimens from one grain per pound of peas upwards, in one case<sup>2</sup> the enormous amount of copper, equal to twenty-six grains of crystallised copper sulphate per pound of peas, was detected. Tschirch would limit the amount to 50 milligrammes per kilo of vegetables. In this proportion he believes it to be absolutely harmless, even if a kilogramme of vegetables thus artificially coloured be eaten by one person daily. The possible toxic effects that may accrue from the consumption of vegetables thus adulterated is repeatedly discussed before the law courts, with varying results— a certain percentage of copper is pronounced injurious on one occasion and not on another. Except in the case of those who from idiosyncrasy are unusually susceptible to the influence of copper, it is very unlikely that poisonous symptoms would manifest themselves after a single meal, of which preserved peas containing a small amount of copper formed a part, although, as stated by Lehmann,<sup>3</sup> as much as 195 milligrammes of copper might be taken with a single meal without the copper being perceptible to the taste. Whilst it is difficult to do other than admit that the occasional use of vegetables thus adulterated would not be attended by symptoms of poisoning, it is at the same time to be regretted that the door is left open for the introduction of a substance which might be productive of harm. Any mischief that resulted would be due to the copper absorbed, and to its local action on the gastric mucous membrane, the vegetables themselves do not appear to be affected so far as their digestibility is concerned. Experiments by Ogier<sup>4</sup> and by Charters and Snodgrass<sup>5</sup> on the artificial digestion of greened vegetables show that the presence in small amount of a salt of copper exercises no adverse influence in this respect. In greened vegetables, copper probably exists as an insoluble leguminate from which the metal is liberated and rendered soluble by the action of the gastric and pancreatic digestion. In Germany, Austria, Belgium, Spain, Russia, and in most

<sup>1</sup> *Das Kupfer vom Standpunkte der gerichtlichen Chemie*, 1893

<sup>2</sup> *Sanitary Record*, 1877

<sup>3</sup> *Munchener med. Wochenschr.*, 1891

<sup>4</sup> *Laboratoire de Toxicologie*, 1891

<sup>5</sup> *The Lancet*, 1892

of the Swiss Cantons, the use of copper for colouring green vegetables is forbidden. In France, where the system of greening vegetables is largely carried out, the law which formerly prohibited the use of copper salts for this purpose has been repealed. In Italy 0.1 grm of metallic copper per kilo of vegetables is allowed. The New York Board of Health allow canned peas to be sold that contain not more than three-quarters of a grain of metallic copper—equal to three grains of the crystallised sulphate—per pound, provided that the label on each tin contains a statement to that effect. In England the law prohibits the addition, to articles of food, of any material which renders the food injurious to health, leaving it to the Court before which the case is tried to decide as to what is, and what is not, injurious to health.

In relation to this subject the question of the natural occurrence of copper in certain vegetables has to be considered, minute quantities have been found in wheat, coffee, potatoes, and a number of other articles, of every-day consumption, including wine, spirits, and effervescing waters. Paul and Cownley<sup>1</sup> found that oysters contain from 1.81 to 3.03 of copper per 10,000 of substance. This, to a certain extent, accounts for the fact that copper is almost invariably present in the human body, of which it has been erroneously assumed to be a physiological constituent. It is probable that copper is being continuously introduced into the system from the source just mentioned, or from the use of copper or brass cooking utensils and hot water apparatus.

When making the investigations on the elimination of lead mentioned in the preceding section, although the patients were not taking any substance known to contain copper, the author rarely failed to obtain evidence of its presence in the *fæces*. The amount varied, it was sometimes considerable, as much as 2 milligrammes of metallic copper being eliminated in the twenty-four hours. Copper was not detected in the urine in these cases. The *fæces* from a number of individuals were subsequently examined for copper, with the result that traces at least, and often much more, were invariably obtained. These analyses show that copper is almost constantly present in the system, and they also prove that the metal is chiefly eliminated by the bowels, corroborative of the latter statement is the fact that when a soluble salt of copper is therapeutically administered, so that one or more grains are taken daily, although minute quantities may be found in the urine, the bulk is eliminated in the *fæces*. It is probable that, like lead, copper is partially retained in the system, but the accumulation takes place more slowly.

**Chemical Analysis.**—Organic matter may be got rid of either by the dry or the moist way. If the former is adopted, the residue, after evaporation of the nitric acid, will probably betray the presence of copper by greenish or bluish tinge.

**Tests.**—The clear solution may be tested with potassium ferrocyanide, which gives a chocolate-brown precipitate, ammonia-water gives an azure blue, potassium sulphocyanide, in dilute solution, gives an emerald green, and in stronger solution an olive green, in either case, on the addition of ammonia-water, the ordinary blue reaction of that reagent with copper is obtained. If a drop of a solution of a copper salt, having a slightly acid reaction, is allowed to remain for a minute or two on the bright blade of a knife it leaves a deposit of metallic copper. Another method is to place in the liquid a bright steel needle, or piece of iron wire, the resulting film of copper may be dissolved in a few drops of ammonia water, to which it imparts a blue tint, this test may be used in the presence of organic matter—tinned peas, for example.

<sup>1</sup> *Pharm Journ*, 1896.

**Quantitative** estimation may be made by precipitating the copper as sulphide, and then dissolving it in strong nitric acid, the acid is evaporated and the residue gradually heated to a full red heat until all the combined nitric acid is driven off. The result is cupric oxide. 100 parts contain 79.85 parts of metallic copper. If only a minute quantity of copper is present, the fluid obtained after destruction of the organic matter is best dealt with by electrolysis, as described in the last section. The deposit of copper is dissolved off the platinum with dilute nitric acid, aided by heat, and, if the amount is very small, estimated volumetrically, if larger, it may be treated with  $H_2S$ , converted into oxide and weighed.

### SILVER.

Acute poisoning by a salt of silver is exceptionally rare, and usually results from the accidental swallowing of a piece of "lunar caustic" which is being used to cauterise the throat.

**Symptoms of Acute Poisoning by Silver Nitrate ( $AgNO_3$ )**—When swallowed in the solid it acts as a violent irritant and corrosive on the mucous membrane of the stomach. Pain is felt in the stomach and abdomen, followed almost immediately by vomiting and probably purging, the early vomited matter consists of cheesy masses of coagulated mucus, which darken on exposure to light, blood may be present both in the vomit and in the dejections. Collapse, cardiac depression, and cramps may occur. Silver is eliminated to some extent by the bowels and kidneys, but most of that which is received into the system is deposited in the metallic state in the tissues.

**Fatal Dose.** Not determined. Scattergood<sup>1</sup> reports a case occurring in an infant, in which a piece of lunar caustic, three-quarters of an inch long, accidentally slipped down the throat, although antidotal treatment was at once resorted to, the child died in six hours. A similar and fatal case happened some years ago to an adult in Manchester.

**Treatment.**—Common salt and water, followed by an emetic, or the stomach-tube, afterwards, white of egg and ice. If the solid lunar-caustic has been swallowed, an emetic is preferable to the stomach-tube. If the mass is not returned it should be removed by laparotomy.

**Post-mortem Appearances.**—Indications of the caustic effect of the poison may be present, as streaks or patches of a greyish-white colour on the parts with which it came in contact. When solid lunar caustic is swallowed, the stomach suffers most severely at the lower part, where the caustic lies. In addition there will be inflammation of the stomach, and probably of the duodenum.

### Chronic Poisoning by Silver.

This usually results from the prolonged internal use of a silver salt medicinally. Instances have occurred in which absorption, resulting in chronic poisoning, has taken place from long-continued application of the nitrate to granulations, workers in metallic silver have also suffered from local symptoms affecting the skin. A common result of chronic poisoning by silver is discoloration of the skin—argyria—due to depositions of particles of the reduced metal in the papillary layer of the corium, not in the rete mucosum, as is the case in physiological skin-pigmentation, on account of the nature of the pigment and its position, the discoloration is very permanent. A dark line is also formed on the margins of the gums, the mesentery, kidneys, and other glandular organs have been found stained. In animals, staining does not take place, but disturbances of nutrition, paralysis and fatty degeneration of the liver and kidneys

<sup>1</sup> *Brit Med Journ*, 1871.



occur Gowers<sup>1</sup> relates the case of a man who, after taking silver medicinally for years, suffered from paralysis of the long extensor of the fingers, and of the extensors of the phalanges of the thumb, on both sides, on the right, the radial extensors of the wrist were also paralysed, argyria and the black line on the gums were present

**Chemical Analysis.**—Destruction of organic matter by the moist method is not feasible, as the silver chloride which would be formed is insoluble Incineration may be adopted, in the course of which, if the poison is present in not too small amount, a coating of metallic silver will be deposited on the bottom and sides of the capsule due to the reducing action of the organic matter

**Tests.**—On adding dilute hydrochloric acid a white curdy precipitate is produced, which is insoluble in nitric acid, but is soluble in ammonia water. A solution of caustic potash gives a brownish precipitate insoluble in excess, but soluble in ammonia and in nitric acid Potassium iodide gives a yellow precipitate, and potassium chromate a red precipitate The quantity of silver may be estimated by precipitating it from a solution of the nitrate by means of sodium chloride, filtering through a tared filter, drying and weighing the precipitate, 100 parts of silver chloride contain 75.28 parts of metallic silver

## ZINC.

Acute poisoning by zinc is limited to two of its salts— the sulphate and the chloride The action of the two salts is different— the sulphate when taken in poisonous doses is simply an irritant, the chloride is a corrosive

### Acute Poisoning by Zinc.

**Zinc Sulphate** ( $\text{ZnSO}_4 \cdot 7\text{H}_2\text{O}$ ), or white vitriol, in naked-eye appearance very closely resembles Epsom salts, for which it has been accidentally administered

The **symptoms** produced by a poisonous dose are violent vomiting, pain in the stomach and abdomen, metallic, astringent taste, which quickly follow the act of swallowing the poison, to these purging usually succeeds The immediate emesis, together with the comparatively feeble toxic action of zinc sulphate, render a fatal issue exceptional, when death takes place it is in consequence of exhaustion

**Fatal Dose.**—Not known Recovery has followed one ounce

**Zinc Chloride** ( $\text{ZnCl}_2 \cdot \text{H}_2\text{O}$ ), known in commerce in the form of Burnett's fluid, and also as soldering fluid, is a violent corrosive

**Symptoms.**—Severe burning sensation in mouth, throat, stomach, and abdomen, followed by immediate vomiting and diarrhoea, with severe tenesmus, and distension of the abdomen, the ejected matters contain traces of mucous membrane and of blood Profound collapse is shown by cold surface, clammy sweat, thin pulse, great prostration, and, in immediately fatal cases, coma, with irregular breathing It is not uncommon for the acute symptoms to mitigate for a time, and then, owing to progressive disorganisation of the tissues of the digestive tract, to recur after an interval of days, or even weeks Ward<sup>2</sup> records an instance where several men on board ship were attacked with symptoms resembling those of cholera, caused by drinking water in which zinc plates were placed, to prevent corrosion of the boiler from whence it was obtained.

<sup>1</sup> *Diseases of the Nervous System*, 1892

<sup>2</sup> *The Lancet*, 1886.

Death has resulted from the application of chloride of zinc paste as an escharotic. The elimination of zinc takes place by the bowels and to a lesser extent by the kidneys.

**Fatal Dose.**—Six grains have proved fatal, but recovery has followed three or four drachms of the solid salt. An infant aged fifteen months recovered after swallowing one teaspoonful of soldering fluid.

**Treatment.**—Poisoning by the **sulphate** will probably require little more than attention to the symptoms, usually the stomach spontaneously relieves itself of the poison, if not, the tube should be used, then warmth should be applied, and stimulants and opium given if required. In poisoning by the **chloride**, potassium or sodium carbonate, tannic acid, white of egg, milk, and demulcents should be administered, followed by opium if necessary.

**Post-mortem Appearances.**—The **sulphate** only gives rise to the appearances usually observed in acute gastro-enteritis, as previously stated, the poison is purely an irritant and, therefore, causes no immediate destruction of tissue. With the **chloride** the case is different. If the patient dies shortly after swallowing the poison, the digestive tract, from the mouth down to the stomach, possibly as far as the duodenum, will show more or less indications of corrosion, there will be patches of softened membrane having a white appearance, which in parts may be detached, together with the usual signs of acute inflammation. If the patient survives the reception of the poison for some weeks, the gastric mucous membrane will probably be completely disorganised and in parts replaced by cicatricial tissue. Jalland<sup>1</sup> records the case of a man who committed suicide by swallowing an unknown quantity of a saturated solution of zinc chloride, he died on the seventy-ninth day. At the necropsy the stomach was found to be completely destroyed, the remains consisted of a sausage-like mass of inflammatory adhesions without any trace of mucous membrane, the cavity—four inches long and three-quarters of an inch in diameter—resembled that of a chronic abscess. A man aged fifty years came under the author's care in hospital after swallowing about one ounce of crude hydrochloric acid "killed" with zinc—that is, with as much zinc in solution as would combine with the acid. This solution of zinc chloride he used at his work. After the immediate effects of the poison had passed off, he improved up to the fourteenth day, when he began vomiting bilious matter which contained sloughs. He progressively wasted, and died from inanition forty-three days after taking the poison. At the autopsy the stomach was seen to be dilated, with very thin walls which had a steel-grey colour, the mucosa was uniformly smooth, the rugæ being absent, and the pyloric orifice was excessively contracted. There were no signs of ulceration.

**Chronic Poisoning** by zinc has been described, chiefly in smelters of the metal. The symptoms to some extent resemble those produced by lead—derangement of the digestive organs, colic with constipation, or, more frequently, diarrhœa, indications of peripheral neuritis have been observed. Gastric symptoms have also been described from drinking water or milk stored in zinc-lined vessels, the zinc used for "galvanising" iron vessels is impure, and fluids containing chlorides will act on it. It is very doubtful whether chronic zinc poisoning actually occurs, and probably the symptoms ascribed to it have really been due to lead, cadmium, or other impurity in the zinc or its compounds.

**Zinc chloride** is sometimes used as "filling" in certain fabrics, and may give rise to acute dermatitis in the wearers of garments made of such fabrics.<sup>2</sup> Recently the author analysed a number of textile fabrics, with respect to the

<sup>1</sup> *Brit. Med. Journ.*, 1887.

<sup>2</sup> *The Lancet*, 1898.

"filling" used, and found that several contained as much as four and five per cent by weight of zinc chloride

**Chemical Analysis**—In neutral or alkaline solution zinc may be precipitated from organic admixture by sulphuretted hydrogen. The precipitated sulphide will probably carry down with it some organic matter, and, therefore, if weighed would indicate an amount in excess of that which was really present, for this reason it is better to convert the sulphide into nitrate or sulphate, and then to precipitate it as carbonate by boiling with sodium carbonate in excess. After being well washed with hot water, the precipitate is strongly ignited so as to convert it into oxide, and it is weighed. 100 parts of zinc oxide contain 80.26 of metallic zinc.

**Tests.**—A white sulphide is formed in neutral or alkaline solutions on the addition of ammonium sulphide, or of sulphuretted hydrogen, the precipitate is insoluble in a solution of potassium hydroxide. A solution of potassium hydroxide added to a solution of a salt of zinc produces an opalescent, gelatinous precipitate, which has a tendency to adhere to the sides of the test-tube, it is soluble in excess. The same result is produced by ammonia water, unless free acid or ammonium salts are present, when no precipitate is formed. Potassium ferro-cyanide gives a pale gelatinous precipitate, and potassium ferricyanide a fawn-coloured precipitate. Ammonium carbonate gives a white precipitate soluble in excess. If zinc oxide is strongly heated it turns yellow, becoming white again on cooling, when moistened with cobalt nitrate and heated with the blowpipe flame, zinc oxide forms a green pigment. Rinman's green.

### CADMIUM.

Chronic poisoning by cadmium occurs occasionally among spelter workers. Stevens<sup>1</sup> believes that cases of cadmium poisoning are often diagnosed as a typical lead poisoning. He describes the case of a spelter worker of 67, who for ten years had been thought to suffer from plumbism, the prominent symptoms being weakness, wasting, and bronchitis. Post-mortem the kidneys showed chronic interstitial inflammation, and the heart was hypertrophied. The liver contained no lead, but 0.91 grain of cadmium per pound, and 0.77 grain of zinc per pound were present. Stevens has seen eight other cases. Chronic cadmium poisoning differs from plumbism in that severe colic is absent, diarrhoea may occur, and there is well-marked epigastric pain or tenderness.

### TIN.

Poisoning by the salts of tin is exceptionally rare, and has only resulted from accident, for the most part from the use of tinned meat and fruit. In Belgium and France it has been found that some confectioners put stannous chloride into gingerbread in order to obtain with inferior materials an appearance like that legitimately due to fine flour, in some instances as much as 5 kilogrammes per 200 kilos of bread were present. After making a series of experiments, Pouchet and Riche<sup>2</sup> have recommended legal prohibition of the practice on the ground of its being injurious to health. No fatal case of poisoning by tin has been yet recorded. A case<sup>3</sup> that has more than once been cited as one of fatal poisoning by the chloride, was really one of hydrochloric acid poisoning, and is so stated by the original reporter, half a teacupful of hydrochloric acid, having some tin in solution, was swallowed, and ultimately caused

<sup>1</sup> *Journal of Industrial Hygiene*, August, 1920

<sup>2</sup> *Annales d'Hygiène*, 1892

<sup>3</sup> *Med Times*, 1841

death Six simultaneously occurring cases of alleged tin poisoning from "canned" tomatoes are recorded by Campbell,<sup>1</sup> five being children and one an adult, one of the children, a girl two years of age, had offensive motions which contained blood, and she died collapsed, the tomatoes were found to contain tin salts, but the toxic effects were probably due to the presence of some decomposition product derived from the fruit

**Symptoms.**—Metallic taste, vomiting and diarrhœa, with pain in the stomach, pain in the head has also been observed In some cases the poison has depressed the action of the heart, Luff<sup>2</sup> records the cases of four adults who suffered from severe symptoms of poisoning after having eaten some tinned cherries, in all four the pulse was feeble, rapid, and irregular, and the surface was cyanotic Luff found 1·9 grains of stannic oxide, derived from the solder used in the construction of the tins, per ounce of the cherry juice, the doses of tin malate respectively received by the sufferers were calculated to range from four to ten grains All the cases recovered Sedwick<sup>3</sup> relates how nine persons, after eating pears which had been stewed in a newly tinned pan, were simultaneously attacked with diarrhœa, vomiting, and abdominal pains, the juice of the fruit was found laden with tin salts Four persons were attacked with griping pains and nausea which, in two, was followed by vomiting one hour after eating tinned rhubarb They all quickly recovered The author found that the rhubarb-juice contained a large amount of tin oxalate which yielded 2·7 grains of stannic oxide per ounce of juice Tin is eliminated by the kidneys and bowels

**Treatment.**—Empty the stomach, and then give demulcents, white of egg, milk, and ice, with opium if necessary

**Chemical Analysis**—Organic matter may be destroyed by the moist method and the resulting solution precipitated with  $H_2S$  The sulphide may then be ignited and converted into stannic oxide, 100 parts of which contain 78·38 parts of metallic tin

**Tests.**—With mercuric chloride and a little hydrochloric acid stannous chloride gives a white precipitate of mercurous chloride, which turns grey and subsequently black (hastened by boiling) from the formation of finely divided metallic mercury With gold chloride it gives a purple precipitate If a little mucine dissolved in a few drops of strong nitric acid and then diluted with about fifty times its volume of water, is boiled, and allowed to cool, a reddish fluid is obtained, a few drops of this added to a solution of a tin salt produces a lilac colour Stannic salts give a yellow precipitate with  $H_2S$ , stannous salts a dark brown, after neutralising with ammonia both sulphides are soluble in ammonium sulphide Both stannous and stannic salts are precipitated by potassium hydroxide and are soluble in excess, and also by ammonia, the precipitates being insoluble in excess

## BISMUTH.

Exceptional cases of poisoning by the subnitrate ( $BiONO_3, H_2O$ ) have occurred both from internal and external use, but instances have become more numerous since the introduction of Beck's method of treating sinuses by the injection of bismuth paste, and the use of bismuth in skiagraphy

**Symptoms.**—Salivation, metallic taste, pain in stomach, vomiting and urging, the dejections having a greyish-black colour, and collapse The gums may be inflamed or even gangrenous and a violet-black line may be present,

<sup>1</sup> *Therap Gaz.*, 1893

<sup>2</sup> *Brit Med Journ.*, 1890

<sup>3</sup> *The Lancet*, 1888.

the breath is very foul. A disagreeable garlic-like odour has been observed from prolonged medicinal use of bismuth, which it is stated is due to impurities in the form of tellurium or of arsenic. In toxic doses the usual symptoms of gastro-enteritis occur. Bismuth is eliminated in the fæces, urine, and saliva, like lead, much passes directly through the bowels without being absorbed.

Warfield<sup>1</sup> has reported the following case.—A girl, aged nine, was admitted into hospital in September, 1911. She had suffered from a psoas abscess which had been twice incised. In November, 1910, about two ounces of Beck's subnitrate paste were injected into the sinus, which promptly closed, and no paste had ever been extracted. Within two weeks a black line was noticed on the margin of the gums, which had persisted, becoming more or less prominent from time to time. In August, 1911, an ulcer formed on the mucous membrane of the right cheek, and later the right side of the tongue ulcerated. The breath was very fetid and many of the teeth decayed. On the margins of both jaws both on the inner and outer sides was a dark violet black line and there was also violet black discoloration along the whole right edge of the tongue. The right cheek was ulcerated. A skiagram of the right lumbar region showed a shadow 10 cm long and 2 to 4 cm broad indicating that almost all the bismuth remained. No bismuth could be found in the urine. Gradual improvement took place, the ulceration healed, but the line on the gums persisted.

**Fatal Dose.**—In one case two drachms caused death in nine days.

**Treatment.**—Evacuate the poison, and then give ice and opium if necessary.

**Post-mortem Appearances.**—Those due to acute gastritis.

**Chemical Analysis.**—Organic matter may be destroyed by the moist method, the bismuth precipitated as sulphide, and afterwards dissolved in concentrated nitric acid, the solution of the nitrate thus obtained is evaporated to dryness and the residue is dissolved in water with the aid of a little nitric acid.

**Tests.**—Potassium hydroxide produces a white precipitate, insoluble in excess, which becomes yellow on boiling. Dilution with water produces a precipitate, which may be distinguished from the precipitate similarly produced with antimony by its insolubility in a solution of tartaric acid. Potassium chromate produces a yellow precipitate which is soluble in nitric acid and insoluble in potassium hydroxide.

**Quantitative analysis** may be conducted by diluting the solution of the nitrate, adding ammonium carbonate and then boiling for some time. The precipitate, after ignition, is weighed. 100 parts of the oxide contain 89.65 of metallic bismuth.

## IRON.

The salts of iron which come under the observation of the toxicologist are the **sulphate** ( $\text{FeSO}_4 \cdot 7\text{H}_2\text{O}$ ), copperas, or green vitriol, and the **chloride** ( $\text{Fe}_2\text{Cl}_6 \cdot 6\text{H}_2\text{O}$ ) in alcoholic solution known as tincture of iron.

**Symptoms.**—Large doses of the sulphate give rise to a metallic taste, pain in the stomach, vomiting, and purging, the dejected matters being black from the formation of ferrous sulphide. The chloride is a more active irritant, in large doses it has been found to act somewhat like hydrochloric acid. Both these salts, but especially the chloride, are not unfrequently administered in poisonous doses for the purpose of procuring abortion, the inefficacy of the proceeding has been already discussed in the section on criminal abortion. Iron is eliminated by the bowels and kidneys.

**Fatal Dose.**—That of the sulphate is unknown. An ounce and a half of the tincture of the chloride has caused death in about five weeks.

<sup>1</sup> *American Journal of the Medical Sciences*, 1912.

**Treatment.**—Evacuate the poison, then give demulcents, ice, and, if necessary, opium

**Post-mortem Appearances.**—If death takes place in the early stage there will be the usual signs of gastritis, with probably some special discoloration of the membrane due to the action of the metal. In the fatal case from the chloride above mentioned, Christison found a thickened, inflamed condition of the stomach towards the pyloric end

**Chemical Analysis**—Examination of the vomited matter or of the contents of the stomach or bowels is alone required, the physiological presence of iron excludes the evidence of the tissues

**Tests.**—Potassium sulphocyanide produces a bright red coloration with a ferric salt, no change with ferrous salts. Potassium ferricyanide gives a brown coloration with ferric salts and a blue precipitate with ferrous salts. Potassium ferrocyanide with ferrous salts produces a whitish precipitate which becomes blue on exposure, with ferric salts a deep blue precipitate of Prussian blue. Potash gives a red brown precipitate with ferric salts and a whitish precipitate with ferrous salts

The amount of iron present in vomited matter may be estimated by converting the metal into a ferric salt, if it is not originally in that form, and then precipitating the oxide with ammonia, igniting and weighing. 100 parts of ferric oxide contain 70 parts metallic iron

## MANGANESE.

Several salts of manganese, such as the chlorides, the sulphate, and the manganese-alum, may give rise to symptoms of poisoning, but, to the toxicologist, the most interesting combination of manganese is —

**Potassium Permanganate** ( $K_2Mn_2O_8$ ).—In strong solution this salt acts as an irritant and, superficially, as a corrosive. It is quickly reduced in the presence of the organic matter it encounters in the stomach, some, however, is absorbed and causes death from heart paralysis

**Symptoms**—Immediately after a poisonous dose is swallowed, pain is experienced from the mouth to the stomach and it rapidly spreads over the abdomen, which becomes tympanitic and very tender on pressure. Uncontrollable vomiting quickly occurs, and there is intense thirst with great difficulty in swallowing. The breathing is difficult and noisy, owing to the action of the poison on the larynx. The tongue and probably the lips and chin are stained black, or dark brown. The heart's action rapidly becomes enfeebled and then ceases. The case is recorded, by Thompson,<sup>1</sup> of a woman who swallowed from fifteen to twenty grammes of potassium permanganate which produced extensive erosion of the pharynx and tumefaction of the glottis so as to necessitate tracheotomy. The lips and tongue were coal black, the corners of the mouth and the chin being also stained. Death from heart-paralysis, took place in five hours. Post mortem, the mouth, pharynx, and epiglottis were found to be corroded, the mucous membrane of the œsophagus and stomach was pallid. The blood was fluid and cherry red in colour. No manganese could be detected in the blood or the urine. Box and Buzzard<sup>2</sup> record the case of a woman, aged forty seven, who swallowed "a handful" of crystals of potassium permanganate in beer. The lips and chin were stained dark brown and the tongue was almost black. When first seen, the pulse was moderately rapid and of fair tension. Very shortly the breathing became slightly stridulous, and she suddenly fell back pulseless and the breathing ceased, death taking place thirty five minutes after the poison was swallowed. Post mortem, the tongue was swollen and almost black at the front. The mucous membrane of the stomach was coated with a black granular powder, and when this was scraped off the membrane was seen to be intensely hyperæmic, as was also that of the duodenum. The corrosive action of the poison was very superficial. The left ventricle of the heart was hypertrophied and firmly contracted. In small doses potassium permanganate has produced severe symptoms of poisoning. A woman, with the intention of committing suicide, swallowed some permanganate

<sup>1</sup> *Petersb. med. Wochenschr.*, 1895.

<sup>2</sup> *The Lancet*, 1899

lotion, the amount swallowed containing about two grains, severe gastric pain, vomiting, and prostration occurred, followed by speedy recovery. Bidwell<sup>1</sup> relates instances where severe symptoms were produced, in one case by two two-grain doses, and in another by a single dose of one grain, taken for amenorrhœa. Hawthorne<sup>2</sup> relates the case of a woman who, to cure amenorrhœa, swallowed in four days a number of pills each of which contained about two grains of potassium permanganate, the total amount taken being twenty two grains. Great pain and tenderness in the abdomen, with excessive thirst and prostration, almost amounting to collapse, ensued. The urine was scanty, the bowels were regular. The symptoms continued until the following day, when improvement occurred, and in a week the patient was quite well.

### CHROMIUM.

The compounds of chromium which are of interest to the toxicologist are chromic acid, potassium dichromate, and lead chromate.

**Chromic Acid ( $\text{CrO}_3$ ).**—A case of poisoning from swallowing about ten ounces of a solution of chromic acid, such as is used for charging zinc-carbon batteries, is recorded by Limbeck.<sup>3</sup> The poison was taken for suicidal purposes, and the patient, when seen two and a half hours after, had violent pain in the abdomen, accompanied by vomiting and purging, which commenced a quarter of an hour after the solution was swallowed, indications of severe collapse were present—cold surface, small frequent pulse, quickened respiration, slight cyanosis visible on the lips, and mental depression. The stomach was well washed out with 18 litres of warm water, and yet half an hour after, about a litre of a dark brown viscid mass was vomited, which contained much chromic acid, this had evidently passed the pylorus and was regurgitated. The mucous membrane of the mouth was not corroded, but in parts it was coloured yellow, the abdomen was distended and tender. The urine was dark brown-red and contained 5 per cent albumin, the dejections from the bowels for the first twenty-four hours were of a peculiar grey-green colour. Chromic acid was found in the vomit, fæces, and urine, in the vomit it was free, and could be separated by simple filtration, but it could not be detected in the fæces and urine until the organic matter was destroyed. The patient recovered in six days.

White<sup>4</sup> records a fatal case from the external use of chromic acid. A young woman suffering from a mass of papillary growth on the external genitals, was treated by a single application to it of about half an ounce of a solution of chromic acid, 100 grains to the ounce, the vagina and the anus were protected by tampons of cotton wool soaked in carbolised oil. Shortly after, the patient experienced pain and thirst, and she vomited, in twenty-four hours she was collapsed—the surface was pale, the pulse rapid, and the extremities were cold, she died twenty-seven hours after the application. Post-mortem examination revealed nothing of moment—the kidneys were passively congested, the capsules peeling easily, the stomach showed some fine ecchymoses. Chemical analysis of the kidneys and liver demonstrated the presence of a salt of chromium. Fowler<sup>5</sup> relates how one or two drops of a saturated solution of chromic acid accidentally swallowed during an application to the throat, produced violent pain in the epigastrium half an hour after, followed by severe vomiting of a green ropy fluid, the patient was collapsed, the face being pale and anxious. Recovery gradually took place. Glycosuria sometimes follows acute chromic acid poisoning.

**Potassium Dichromate ( $\text{K}_2\text{Cr}_2\text{O}_7$ )** is used for a variety of trade purposes,

<sup>1</sup> *Boston Med. and Surg. Journ.*, 1886.

<sup>2</sup> *Prager med. Wochenschr.*, 1887.

<sup>3</sup> *The Lancet*, 1899.

<sup>4</sup> *University Med. Mag.*, 1889.

<sup>5</sup> *Brit. Med. Journ.*, 1889.

such as staining wood, and for dyeing, and in this way is very accessible to the public

**Symptoms.** A bitter acrid taste, followed by burning pain in the stomach, vomiting, purging, intense thirst, and prostration are usually present, the vomited matter and the motions may contain blood, there is a tendency for the respiration to be affected. Stewart<sup>1</sup> records a fatal case in which a woman became unconscious five minutes after swallowing one ounce of potassium dichromate dissolved in water, there was severe vomiting and purging, collapse, small, thin, irregular pulse, slow, irregular, gasping respiration, with intervals amounting to fifteen seconds between the breaths, the heart continued to beat fully one minute and three-quarters after respiration had ceased. In another case recorded by Turnbull,<sup>2</sup> in which recovery took place, the respirations were at one time forty-eight to the minute. In all cases the symptoms of collapse and of acute gastritis are very pronounced. Pander<sup>3</sup> found experimentally that chromium salts produce disturbance of the respiration and of the central nervous system, the heart's action not being directly affected. Parenchymatous nephritis and blood-changes occur in chronic cases. Chromium is eliminated chiefly by the bowels, to a lesser extent by the kidneys.

In the manufacture of potassium dichromate the workmen are liable to suffer from sores or "chrome holes" on the hands and face, which have an indurated, cup-like border, and strongly resemble hard chancres. Destruction of the cartilage of the nose also is not uncommon, the septum being perforated at the lower part.

**Fatal Dose.**—Two drachms have caused death in four hours. Recovery ensued after a dose estimated at 273 grains.

**Treatment.**—Stomach-tube, or an emetic, followed by magnesium carbonate or chalk suspended in water or milk. Opium and the usual treatment for excessive vomiting and collapse will probably be required. After free lavage of the stomach with warm water, von Jaksche recommends further washing out with a weak solution of silver nitrate.

**Post-mortem Appearances.**—When death quickly follows the reception of the poison the mucous membrane of the stomach, presenting the usual appearances of acute inflammation, has been found to be superficially eroded in parts, in addition, it has been observed to be stained a deep olive-green, due to the presence of oxide of chromium, green-stained mucus has been found in the stomach. Externally, yellow stains may be present on the lips and the corners of the mouth. Ruttan and Lafleur<sup>4</sup> found the blood chocolate-coloured and obtained from it the spectrum of methæmoglobin.

**Lead Chromate** ( $\text{PbCrO}_4$ ) is used as a pigment under the name of chrome yellow.

The symptoms produced by this poison are a complex of those due to chromic acid and those due to lead. Although the salt is insoluble in water, when swallowed in large quantities it may produce symptoms like those of potassium dichromate, it is not nearly so actively poisonous as the dichromate, but it gives rise to after-effects due to the lead which render it a very dangerous preparation. Most of the recorded cases resulted from the use in confectionery of lead chromate as a pigment, or from inhalation of the salt in the process of its manufacture. Stewart<sup>5</sup> gives a clinical analysis of 64 cases of poisoning by chrome yellow, which had been used to colour cakes, the doses were small,

<sup>1</sup> *Brit Med Journ.*, 1888

<sup>2</sup> *The Lancet*, 1892

<sup>3</sup> *Beitrage zur Chromwirkung*, 1887.

<sup>4</sup> *Montreal Med Journ.*, 1888.

<sup>5</sup> *Medical News*, 1887.



but were repeated, the symptoms were those due to lead. In larger doses death has resulted after a period of drowsiness and apathy.

**Chemical Analysis**—After being liberated from organic matter by the moist method, the solution of chromium chloride, with the addition of a little sulphuric acid, is boiled for some time with alcohol until it turns a green colour. The alcohol is then driven off, and ammonia is added in excess, the liquid is again boiled for some time, after which the precipitate of chromium oxide is filtered off, dried, and ignited. 100 parts contain 68.62 of chromium.

**Tests.**—The soluble salts of chromic acid in acid solution are changed to green by sulphuretted hydrogen. Barium chloride gives a yellow precipitate soluble in hydrochloric acid. Silver nitrate gives a crimson precipitate soluble both in ammonia and nitric acid. Lead acetate gives a yellow precipitate soluble in potassium hydroxide, and with difficulty in nitric acid. With the aid of heat, sulphuric acid and alcohol reduce chromates to chromic oxide salts, changing the colour from yellow, or red, to bluish-green. Lead chromate digested with dilute sulphuric acid yields a precipitate of lead sulphate, chromic acid remaining in solution, if the solution is filtered, and to the filtrate a little hydrochloric acid is added, and then sulphuretted hydrogen is passed through it, the colour after a time changes from red to green, the precipitate of lead sulphate may be reduced with the blowpipe to the metallic state, converted into a nitrate, and subsequently tested.

## NICKEL.

The poisonous effects of this metal have been chiefly produced by the tetracarbonyl,  $\text{Ni}(\text{CO})_4$ . Nickel carbonyl is a highly refractive liquid which becomes gaseous at  $104^\circ \text{F}$ . In human beings, its toxic action has been produced by the inhalation of air contaminated with its vapour. The symptoms produced comprise headache, dyspnoea with rapid respiration, lowering of the temperature and, in fatal cases, convulsions and coma. From experiments on animals, M'Kendrick and Snodgrass<sup>1</sup> state that the toxic effects of nickel carbonyl resemble those produced by carbon monoxide, carboxyhaemoglobin being found in the blood. Herriot and Richet<sup>2</sup> come to a like conclusion. Vahlen<sup>3</sup> believes that they are not produced by the carbon monoxide which is split off, nor by the metal, but by the specific action of the compound, nickel carbonyl. Mittasch<sup>4</sup> noted the rapid breathing and dyspnoea in animals. In 1902, three men were fatally poisoned by this carbonyl at a nickel chemical works, and several others were made ill, but they recovered. The symptoms were headache, vertigo, fever, and rapid breathing. In the fatal cases, the lungs were found to be congested and oedematous; the brain was also congested.

## GOLD AND PLATINUM.

Poisoning by the salts of these metals is rare, but as they are used in photography, which of late has become a fashionable amusement, their toxic effects are likely to be more frequently encountered in the future, as the two following cases indicate.—

Stevenson<sup>5</sup> relates the case of a boy who swallowed part of the contents—something under 12 grains—of a photographer's tube of gold trichloride which a playmate had found in a dust-heap; he vomited violently and became collapsed, the lips, tongue, teeth, and inner side of the cheek were stained purplish-black by the reduced gold. Diarrhoea followed, accompanied by tenderness in the epigastrium with retracted abdomen, both vomit and motions were free from blood. Gold was found in the early vomit and in the faeces, but not in the urine. The treatment was symptomatic, and recovery speedily took place.

Hardman and Wright<sup>6</sup> relate an interesting case in which a woman went to a chemist's

<sup>1</sup> *Brit. Med. Journ.*, 1891.

<sup>2</sup> *Compt Rend Soc Biol.*, 1891.

<sup>3</sup> *Arch. j. exp. Pathol.*, 1902.

<sup>4</sup> *Ibid.*, 1903.

<sup>5</sup> *Guy's Hosp Reps.*, 1893.

<sup>6</sup> *Brit. Med. Journ.*, 1896.  
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shop to buy a teething powder for her seven-months old infant, by mistake she picked up from the counter a small packet containing 8 grains of potassium chloro-platinite, which had just been weighed out for a photographer, and gave it to the child. Vomiting and diarrhoea occurred, with the usual symptoms of gastro-enteritis, the child became collapsed and, in spite of treatment, death from cardiac failure took place in five hours. At the autopsy the mucous membrane of the stomach was found to be pale except for a patch of brownish yellow staining on the posterior wall, the spleen was enlarged and the kidneys, highly congested, displayed punctiform hæmorrhages, a chronic intussusception was found which possibly had something to do with the fatal termination. Platinum was found in the stomach and intestines.

**Chemical Analysis.**—Any organic matter may be destroyed either by the moist way or by heat, and an auric or platinum chloride obtained.

**Tests.**— $\text{AuCl}_3$  forms with water a strongly acid, yellowish solution, if to it a mixture of stannous chloride with a little stannic chloride be added, the purple of Cassius is produced. Oxalic acid reduces  $\text{AuCl}_3$ .  $\text{PtCl}_3$  with  $\text{KOH}$ , or  $\text{NH}_4\text{OH}$ , gives a yellow crystalline precipitate. Oxalic acid does not reduce platinum salts.

## CHAPTER XXXI

### NON-METALLIC ELEMENTS.

#### PHOSPHORUS.

THE toxic properties of phosphorus are chiefly utilised as a means of committing suicide, to this end some kind of vermin-killer or rat-paste containing phosphorus is swallowed. These pastes consist of fatty matter, amongst which finely divided phosphorus is distributed to the extent of about 3 or 4 per cent mixed with flour, sugar, and usually some pigment, a small-sized pot will contain 4 to 6 grains of phosphorus. In default of the paste, the heads of a number of matches are sometimes detached, mixed with water, and swallowed, children are occasionally accidentally poisoned by sucking off and swallowing the heads of matches. Only the matches made with yellow phosphorus are poisonous, "safety-matches" which solely ignite on surfaces of red amorphous phosphorus are inert. In 1919, seven deaths (one accidental, six suicidal) were recorded from phosphorus poisoning, in England and Wales.

When phosphorus is swallowed, especially when the poison is in a finely divided state (in which it invariably exists in the combinations used for suicidal purposes), it is capable of being absorbed as such, without previously undergoing oxidation.

#### Acute Phosphorus Poisoning.

**Symptoms.**—After a poisonous dose of phosphorus is swallowed, pain in the stomach, followed by vomiting, comes on in from a few minutes to twelve or twenty-four hours. In exceptional cases the symptoms have first appeared at a still more remote period, not even until the second or third day, usually they occur in from two to five hours. In a dark place the matters first vomited and the patient's breath often appear luminous. A phosphorus- or garlic-like odour may be perceived in the breath both by the bystanders and by the patient himself, after the stomach has been well emptied, the subsequent vomited matter will not be phosphorescent, although the odour may for a time persist.

Intense thirst, eructations, and a burning sensation in the throat and stomach are experienced. Diarrhœa is more frequently absent than present, it occurs in about 25 or 30 per cent of cases. In *rapidly fatal cases* the collapse which accompanies these symptoms increases, the vomiting continues, the rejected matter probably contains blood, the abdomen is distended and is exceedingly tender, the patient is anxious, restless, and exhausted, and death takes place in eight or ten hours. Delirium and convulsions may precede death.

Such, however, is *not the usual course* of acute phosphorus poisoning, in most cases the *primary symptoms diminish* in intensity, sometimes so much so as to convey to the inexperienced observer an impression that the danger is past. The stage of partial passivity may last for two or three days or even longer, the patient appearing to have little the matter with him, or he may continue to have occasional attacks of vomiting, a distended, tender abdomen, and a quick feeble pulse. The diarrhœa, if present in the initial stage, may cease and give place to pronounced constipation. The tongue is coated and the thirst continues. In exceptional instances two or even three weeks have elapsed before the appearance of the secondary symptoms. West<sup>1</sup> records a case in which after a piece of phosphorus-paste the size of a walnut was swallowed there were no symptoms for six weeks, the patient then commenced to be ill, and died in six days.

The commencement of the **secondary symptoms** is often indicated by yellow discoloration of the sclera, the skin usually participating in the icteric hue, which may spread over the whole body, pain is felt in the epigastric region, and on examination the liver is generally found enlarged, sometimes the spleen is also enlarged. The abdomen becomes greatly distended and tympanitic, vomiting returns, and the bowels are more or less relaxed, both the vomit and the motions containing much blood. A general hæmorrhagic tendency is evinced by bleeding from the nose, and in women from the vagina, and by the formation of purpuric spots and ecchymoses under the skin and mucous surfaces. The urine becomes high-coloured and scanty, it has a strong acid reaction, and frequently contains bile-pigments, albumin and blood-colouring matter. The pulse is quickened—80 to 100 per minute—the temperature varies, but usually is not much above the normal. Pain in the head, restlessness, and sleeplessness occur, together with affections of the special senses—as singing in the ears, deafness, and impaired vision. In some instances formication and cramps have been observed. Bollinger<sup>1</sup> relates a case in which a girl died in four and a half days after swallowing the heads of some lucifer matches, paresis of the extremities occurred on the third day, and on the fourth the feet were completely paralysed. On section the spinal meninges, especially at the nerve-roots in the dorsal and lumbar regions, were infiltrated with blood. In fatal cases the condition of the patient rapidly becomes worse, the pulse is irregular, and a state of stupor or coma supervenes, which is soon followed by death. The mind may remain clear until nearly the last, or there may be acute delirium, with or without convulsions. The temperature sometimes sinks considerably towards the end, in other cases it rises, and has been known to continue to rise for a short time after death. Exceptionally a diminution in the size of the liver has been noticed during the last days of life. In a few cases recovery has taken place after the occurrence of enlargement of the liver and jaundice, the liver gradually acquiring its normal size, and the skin its usual colour.

**Fatal Dose.**—The smallest fatal dose was probably  $1\frac{1}{2}$  grains. Recovery

<sup>1</sup> *The Lancet*, 1893.

<sup>1</sup> *Deutsch. Arch. f. klin. Med.*, 1869.

has taken place after 4 to 6 grains, swallowed in the form of rat-paste. Death has occurred in twelve hours, and even earlier; more frequently it is delayed until the second to the fourth day, in the greater number of cases it takes place within a week, although life has been prolonged for over twice that period.

**Treatment.**—Evacuate the contents of the stomach with the stomach-tube or give doses of two or three grains of copper sulphate dissolved in water. The copper salt acts as an emetic, some of it, reduced by the particles of phosphorus is precipitated on them as metallic copper, and thus tends to render them inert. Half-drachm doses of old unrectified turpentine, especially the French variety is recommended as an antidote, but it is difficult to obtain. If, as is supposed by some, the efficacy of the turpentine depends on its containing oxygen in the form of ozone, it is probable that "Sanitas," which consists of artificially oxidised turpentine, containing hydrogen peroxide, might be equally or even more efficacious. It has been recently recommended, in cases of acute phosphorus poisoning, to wash out the stomach with a 0.1 per cent aqueous solution of potassium permanganate. Purgatives should afterwards be given and theriac, demulcents, and morphine. The fæces voided a week or longer after the poison was taken have been observed to be luminous, showing that it persisted in an active form, repeated evacuation of the bowels, therefore, should be procured, but not with castor oil, no oily nor fatty matter should be given as such substances are solvents of phosphorus, and would promote absorption. The inhalation of oxygen, especially along with ozone, is sometimes beneficial.

**Post-mortem Appearances.**—The cavities of the body, when opened, may yield the odour of phosphorus, and in some cases phosphorescence has been observed. The œsophagus is not usually inflamed. The mucous membrane of the stomach is yellowish- or greyish-white, it is infiltrated, and the epithelium is cloudy and swollen, limited erosions and ecchymoses may be present. The epithelial cells of the gastric glands show more or less advanced fatty degeneration, at first they are filled with fine granular matter, later on fat globules appear. Similar changes may be found in the duodenum, which, as well as the stomach, may contain blood-stained fluid. The intestines frequently show no changes other than small ecchymoses, in some instances they have been found inflamed. The heart and the kidneys show signs of fatty degeneration. The spleen is usually enlarged. Elkins and Middlemass<sup>1</sup> found fatty changes in the nerve cells of the cerebral cortex.

The most marked appearance is that presented by the **liver**; in the greater number of cases the organ is considerably enlarged, but it may be unaltered in size, and has been found contracted. It is of a doughy consistence, easily torn, varying in colour from a bright to a dirty pale yellow, the entire surface may be uniformly tinged, or it may present a marbled appearance—the normal hue of the liver being preserved in parts, this is more especially observable on making a section of the organ. Small hæmorrhagic spots are often present on the surface and in the substance of the liver. Microscopical examination shows that the normal appearance is due to the presence of a large amount of fat. It was formerly disputed whether the fat resulted from degeneration of the liver cells, or simply from infiltration, recent investigations have shown that it is the result of metamorphosis of the tissues it replaces. Nathanson<sup>2</sup> extracted the fat out of hardened pieces of liver by boiling them in ether. In the normal liver no change was produced, in fatty infiltration the fat was removed, leaving the liver-cells intact, in phosphorus-liver the degenerated tissue was dissolved out by the ether, and consequently the liver structure

<sup>1</sup> *Brit. Med. Journ.*, 1891.

<sup>2</sup> *Dissert.*, 1890.

was obliterated Endeavours have been made to trace the steps of fat formation by ascertaining the amount of lecithin in the phosphorus-liver, and comparing it with that present in the normal liver Stolkow<sup>1</sup> found that phosphorus causes an increase in nuclein, which leads to excess of lecithin, from which the phosphorus is split and the fat formed Leo<sup>2</sup> found no increase Heffter<sup>3</sup> found a distinct diminution in lecithin, averaging nearly 50 per cent, the greater the amount of fat in the liver, the less the amount of lecithin He believes it unlikely that phosphorus causes fatty degeneration of albuminoid substances, with the formation of lecithin as an intermediate product, more probably the store of lecithin already formed in the liver-cells undergoes chemical change The processes by which the liver-changes are produced are still undetermined, by some they are referred to the direct action of the phosphorus on the cells, and by others to morbid processes set up in the parenchyma the jaundice probably results from blocking of the primary gall ducts by tumefied epithelium The fœtus of a woman who died from acute phosphorus poisoning shortly after delivery, afforded evidence of the action of the poison in the form of fatty degeneration of the liver-cells, together with numerous ecchymoses in various organs

The resemblance between acute phosphorus poisoning and acute yellow atrophy of the liver has long been observed, though there are obvious differences the phosphorus liver is usually hypertrophic with well marked acini, in acute yellow atrophy the liver is small, and all indications of acini have disappeared It has been asserted that the morbid anatomy of the acute atrophic liver is characterised by inflammation of the intralobular connective tissue, with molecular degeneration of the liver cells, that of the phosphorus liver by simple fatty infiltration of the liver-cells The difference seems to be one of degree Wysz<sup>4</sup> treated some sections of phosphorus liver with turpentine, to remove the fat, and found in some parts that the liver cells were sharply defined, while in others they had entirely disappeared In acute atrophy the morbid processes in the liver appear to progress more quickly than in phosphorus poisoning, when the phosphorus liver has had time to undergo advanced changes, it cannot be distinguished from the acute atrophic liver Heffter states that in chemical composition the two livers are identical In exceptional instances Reiss found proliferation of cells of the interstitial tissue as in acute atrophy Hessler<sup>5</sup> states that out of sixty four cases of phosphorus poisoning, in thirteen the liver was small, as in acute atrophy Heddench<sup>6</sup> reports a well marked case of a girl aged eighteen, who put a box of matches into a gill of warm water for ten minutes and then drank the water jaundice and other indications of phosphorus poisoning supervened, but the liver was diminished in volume from the first The question is not yet determined, but the analogies between acute atrophy and phosphorus poisoning render it probable that both are due to toxic influences

The investigations of Silbermann,<sup>7</sup> Badt,<sup>8</sup> and others show that in acute phosphorus poisoning the blood corpuscles acquire a tendency to agglutinate together, and thus give rise to numerous thromboses in a case recorded by Haberd,<sup>9</sup> the skin of the feet became gangrenous from thrombosis of the veins Jaksch<sup>10</sup> states the alkalinity of the blood is diminished and the number of red corpuscles increased The localised hæmorrhages, so universal in phosphorus poisoning, are probably due to fatty degeneration of the vascular walls, together with the formation of thrombi within the lumina of the smaller vessels

Acute phosphorus poisoning gives rise to certain well-marked alterations in metabolism, the characteristic features of which are rapid cleavage of proteid and excretion in the urine of many of the primary constituents of the proteid molecule in an unchanged state In many cases of acute phosphorus poisoning, the daily excretion of urea has been found to be below the normal, some of the nitrogen thus wanting being represented by an increased

<sup>1</sup> *Arch. f. Anat. u. Phys.*, 1887.

<sup>2</sup> *Zeitschr. f. physiol. Chemie*, 1885

<sup>3</sup> *Arch. f. exper. Pathol.*, 1891.

<sup>4</sup> *Virchow's Arch.*, 1865.

<sup>5</sup> *Vierteiljahrsschr. f. ger. Med.*, Bd 36

<sup>6</sup> *Munchener med. Wochenschr.*, 1895

<sup>7</sup> *Virchow's Arch.*, 1889

<sup>8</sup> *Stoffwechsel bei Phosphorvergiftung* (Diss.), 1891

<sup>9</sup> *Versammlung deutscher Naturforscher u. Aerzte*, 1894

<sup>10</sup> *Deutsch. med. Wochenschr.*, 1893.

percentage in the form of intermediate products, especially ammonia, tyrosin, leucin, and some other amino acids may also be present in the urine. In several instances the total amount of nitrogen excreted daily has been found not to be much below the normal until shortly before death, some observers state that the total nitrogen is diminished. Munzer<sup>1</sup> states that for the first day or two after a poisonous dose of phosphorus, the amount of N excreted is very small, in consequence of the patient being in a state of starvation—taking no food and being unable to retain the fluids he drinks, on the second or third day there is a marked increase in the amount of N excreted, owing to the destructive action of the phosphorus on the tissue proteid, which also leads to an extraordinary increase, for two or three days, of phosphates in the urine. Ammonia has been found in the urine in excess proportional to the absence of urea. Engelen<sup>2</sup> in experiments on animals found only a slight increase. Starling and Hopkins<sup>3</sup> found a large increase in a fatal case of acute phosphorus poisoning, the N represented by the ammonia in the urine being to that represented by urea as 1 to 7, the normal proportion being 1 to 70. Badt in one case found double the normal amount, in a second case 25.8 per cent—more than one fourth of the total N present in the urine. When the amount of urea in the urine represents less than 85 to 90 per cent of the whole of the N present, disease of the liver is indicated, but Munzer,<sup>4</sup> who found that the urea N is diminished to from 70 to 80 per cent, the ammonia N being increased to from 10 to 18 per cent, attributes the increase to the necessity for neutralisation of the acid products developed in excess by the abnormal metabolism, which the ordinary resources of the organism are unable to meet. The deficiency is made good by the utilisation of some of the ammonia derived from the splitting up of the proteid, the ammonia salts thus formed are excreted in the urine. In rabbits poisoned by phosphorus, no increased excretion of ammonia occurs, because their food, being exclusively vegetable, furnishes a sufficiency of alkali to neutralise the acid, so that an ammonia supplement is not required. Some of the excess of ammonia excretion in the human subject may be due to incapacity of the liver to convert the whole of it into urea. Tyrosin has only occasionally been met with, and leucin still more rarely. Poore<sup>5</sup> investigated a case in which a large quantity of phosphorus (8 to 10 grains) in the form of rat paste was taken, jaundice set in on the third day, and the patient died on the fifth day. In some urine obtained during the last hours of life a small amount of tyrosin, but no leucin, was found. Riess<sup>6</sup> found tyrosin in six out of thirty six cases. A few other observers have recorded instances in which tyrosin was present. Leucin has been found by Ossikowski,<sup>7</sup> Blenderman,<sup>8</sup> Rothammer<sup>9</sup> (along with tyrosin), and three or four others. It is to be noted that in several of these cases, the urine in which tyrosin and leucin were found was voided immediately before death. In the urine of rabbits poisoned with phosphorus, Abderhalden and Bergell<sup>10</sup> found glycochol and some other monamino acids. Albumin is commonly present, but not in large amount. Peptones, or substances giving the same reaction, and oxyaromatic acids, such as oxy-mandelic acid, have been found in the urine by one or two observers. V. Noorden states that the oxyaromatic acids are derived from tyrosin, their presence, therefore, is an indication that the oxidation of the aromatic nucleus has been commenced, but is not capable of being carried further. Consequently if a considerable amount of aromatic oxy acids be present in the urine little or no tyrosin is found. Robitschek<sup>11</sup> found abundance of peptones, along with albumin, in the urine of a girl aged nineteen, who died six days after swallowing the heads of five boxfuls of matches, the peptones afterwards diminished in quantity and were entirely absent on the day before death. On the other hand, Marxner states that the peptonuria increases with the severity of the symptoms. Free fat has also been met with.

Defective metabolism is further shown by the presence of non-nitrogenous abnormal bodies in the urine. Sarcosolactic acid has been frequently found. Riess found it in twenty-six out of twenty seven cases. Poore found it in the case previously mentioned. Under normal conditions sarcosolactic acid is excreted as carbon dioxide and water, its presence in the urine, therefore, is very significant of imperfect oxidation. Sugar has been found in the urine in a few cases, in the one recorded by Poore it was present. In Bollinger's case a small quantity was present. Grose<sup>12</sup> found it on the third and fourth day in a child three and a half years old who was fatally poisoned with rat-paste.

In experiments on animals Baur<sup>13</sup> found that the intake of oxygen and the output of

<sup>1</sup> *Deutsch. Arch f. klin. Med.*, 1894.

<sup>2</sup> *Dissert.*, 1888.

<sup>3</sup> *Guy's Hosp. Reps.*, 1890.

<sup>4</sup> *Centralb. f. klin. Med.*, 1892.

<sup>5</sup> *The Lancet*, 1888.

<sup>6</sup> *Real-Encyclopadie*, 1888.

<sup>7</sup> *Wiener med. Wochenschr.*, 1881.

<sup>8</sup> *Zeitschr. f. physiol. Chemie*, 1882.

<sup>9</sup> *Dissert.*, 1890.

<sup>10</sup> *Zeitschr. f. physiol. Chemie*, 1903.

<sup>11</sup> *Deutsche med. Wochenschr.*, 1893.

<sup>12</sup> *The Lancet*, 1889.

<sup>13</sup> *Zeitschr. f. Biologie*, VII. u. XIV.

carbon dioxide are both considerably diminished, thus, taken in conjunction with the increased splitting up of the albuminoids, to a great extent accounts for the accumulation of fat in the various tissues—the non nitrogenous bodies, such as fats, which result from the splitting up of the albumin, are retained in the system, instead of being burnt up into  $\text{CO}_2$  and  $\text{H}_2\text{O}$ . The mode in which phosphorus produces these pronounced alterations in tissue metamorphosis is not known. Typical effects have followed such small doses that it is impossible to refer them to chemical processes directly due to the reducing properties of phosphorus. Playec<sup>1</sup> contends that the absorbed phosphorus does not exist in the free state, and he instances in support of this view the insignificant influence produced by the inhalation of oxygen and ozone on the progress of a case of phosphorus poisoning. He then submits that the absorbed phosphorus is either oxidised or enters into protoplasmic combination, and accepts the latter view. The rapidity of this combination is directly proportional to the amount of oxyhaemoglobin that is in the blood. When free phosphorus is found in the blood or in the tissues, it is in consequence of the ingestion of an excessively large dose. There is no doubt, however, that free phosphorus is absorbed, and, at any rate occasionally, is excreted as such. Whether the presence of phosphorus in the organism determines the change in tissue metamorphosis somewhat after the mode in which a ferment acts, or in some other way damages the physiological properties of cell protoplasm, remains an open question. It seems as though the poison lowers the oxidising capacity of the cells throughout the organism, and to an excessive degree in certain tissues. Jacoby<sup>2</sup> states that there are grounds for believing that changes in the fermentive processes in the organism play an important part in the pathology of phosphorus poisoning.

Large doses of phosphorus, of which some has undoubtedly been absorbed, do not invariably produce the chain of symptoms described. Stevenson<sup>3</sup> records the case of a woman aged twenty-two who swallowed nearly half an ounce of rat-paste. The vomited matters and the breath were luminous, and gave off the odour of phosphorus, the abdomen was distended and tender, severe collapse occurred, but there was no jaundice, the woman completely recovered in five or six days. The author once saw a case under the care of a colleague, Dr Edge, in which more than half an ounce of rat-paste was swallowed by a woman, who subsequently vomited large quantities of blood, the abdomen was very tympanitic and tender, and remained so for weeks, with recurrent hæmatemesis. The urine first passed after the poison was taken smelt strongly of phosphorus, and showed its presence by Mitscherlich's test, demonstrating that some of the phosphorus had been absorbed and excreted unchanged. The woman recovered without the least sign of jaundice, or of any of the symptoms due to changes in metabolism.

**Phosphoretted Hydrogen ( $\text{PH}_3$ )** is a very poisonous gas, it has caused death by being breathed in an atmosphere contaminated with 0.25 per cent. Several fatal cases of poisoning by this gas have occurred on board ships laden with **ferro-silicon**, an alloy of iron and silicon employed in the manufacture of steel. Ferro-silicon is chiefly prepared on the continent and is shipped to this country, where it is transported inland in barges. It contains as an impurity calcium phosphide, from which, under the action of moisture or moist air, phosphoretted hydrogen is evolved. Calcium arsenide is also present, and from this is given off arsenuretted hydrogen. Copeman, Bennett, and Hake<sup>4</sup> record a number of cases of wholesale poisoning among the passengers and crews of vessels carrying ferro-silicon. In 1905, fifty of the steerage passengers of the *Vaderland* from Antwerp to New York were made seriously ill by the fumes given off from the cargo, and eleven died, the deaths at first being certified as due to pneumonia. In 1908, five Russian immigrants were fatally poisoned during the twenty-four hours' voyage of the *Ashlon* from Antwerp to Grimsby. Cases in canal boats have been attributed to ptomaine poisoning. The Board of Trade now requires

<sup>1</sup> Pfäuger's Arch., 1904.

<sup>2</sup> The Lancet, 1880.

<sup>3</sup> Zeitschr. f. physiol. Chem., 1900.

<sup>4</sup> Supplement to Report of Loc. Govt Bd., 1908-09.

precautionary measures to be adopted in the transport and storage of this dangerous substance

### Chronic Phosphorus Poisoning.

Ordinary yellow phosphorus is largely used in the preparation of lucifer matches, the manufacture of which formerly gave rise to numerous cases of chronic poisoning. The employment of red phosphorus has done something to abolish the evils which existed in the older match manufactories, but improved hygienic surroundings and greater precautions have done much more to render the occurrence of chronic phosphorus poisoning less frequent.

Chronic phosphorus poisoning is caused by frequent inhalation of the fumes of phosphorus, the special pathological condition produced being necrosis of the bones of the upper and lower jaw, especially of the latter. Phosphorus vapour acts locally on bone tissue wherever the periosteum is exposed, as long as it is covered with mucous membrane it possesses complete immunity. The usual path by which the vapour finds its way to the bone is that afforded by a decayed tooth, or in an interspace where a tooth is absent. The surrounding gum inflames, swells, and separates from the alveolar process, the teeth loosen, and either fall out or have to be removed on account of the pain. The bone is first attacked with periostitis, this leads to necrosis, which in severe cases spreads far beyond the spot originally attacked. The general health suffers, partly from the action of the phosphorus on the system at large, and partly from disordered digestion resulting from imperfect mastication of food, and from some of the pus derived from the diseased jaw finding its way into the stomach. Bronchial catarrh and constipation have been observed to precede the bone necrosis. Stockman<sup>1</sup> regards the necrosis as being due to the action of the tubercle bacillus: the phosphorus fumes erode the bone and weaken its nutrition, and so make it susceptible to infection by the bacilli. According to Arnaud,<sup>2</sup> the phosphorus which is absorbed by workmen in lucifer-match manufactories is, for the most part, slowly eliminated in the urine, which may yield the odour of phosphorus, a slight degree of albuminuria often occurs at the time, which, however, indicates no perceptible pathological disturbance. In healthy subjects, and apart from its action on exposed bone, the phosphorus present in the atmosphere of a match-manufactory does not act as a toxic agent.

The treatment of the bone disease is surgical. As preventives, free ventilation and the employment only of workmen with sound teeth, together with periodic examination of their mouths, are necessary.

**Chemical Analysis.**—Vomited matters, and the contents of the stomach obtained post-mortem, should be examined in the dark for luminous particles, if much phosphorus is present the whole mass will give off phosphorescent vapour. By daylight the substance should be searched for particles of Prussian blue or other pigments with which the phosphorus may have been mixed. The odour of phosphorus is perceptible when present in very small amount, provided that it is not overpowered by the odour of other volatile bodies.

**Tests.**—The most delicate test for phosphorus when in organic admixture is afforded by distillation in the dark, this method, known as **Mitscherlich's test**, is carried out as follows.—The suspected substance if alkaline is acidulated with a few drops of sulphuric acid, and if necessary water added until it is of a fluid consistence, it is then put into a distillation-flask furnished with a condenser, the free end of which dips into a receiver containing a solution of

<sup>1</sup> *Brit. Med. Journ.*, 1899

<sup>2</sup> *Annales d'Hygiène*, 1896



silver nitrate, the condenser is enclosed in a box, the interior of which is painted dead black, and two eye-holes are provided so that the tube may be observed in perfect darkness. Heat is applied to the flask, and if phosphorus is present, even only in minute quantity, the inner tube of the condenser becomes partially or wholly luminous. Certain substances, of which those most likely to be encountered are turpentine (given as an antidote), alcohol, ammonia, ether, and sulphuretted hydrogen, prevent the development of luminosity. A small amount of phenol diminishes the delicacy of the test. The solution of silver nitrate is blackened by reduction of the silver to the metallic state, and it contains phosphoric acid.

Another test is that known as **Dussart-Blondlot's**. Hydrogen is passed through a flask containing the suspected substance, when, if phosphorus is present, some of it combines with the hydrogen, forming phosphoretted hydrogen, which burns with a characteristic flame. The apparatus specially constructed for the purpose enables two flames to be placed side by side for comparative observation—one derived from ignition of the hydrogen before it has passed through the flask containing the suspected substance, and the other after it has done so, both jets must be of platinum, as those made of glass tubing (on account of the soda that is present) do not yield sufficiently non luminous flames. The flame of phosphoretted hydrogen has a green tint in the centre, most marked when it plays on a cool surface. Examined with a spectroscope it gives three lines in the green—one at the E line, another between F and F', and the third between D and E, there are other lines present when the flame is viewed under the most favourable circumstances, but those enumerated are sufficiently distinctive. Hypophosphites gave the same results. A third test, known as **Scherer's**, is founded on the reducing power of phosphorus, by which filter paper moistened with a solution of silver nitrate is blackened if exposed to the action of phosphorus or of phosphorous oxide vapour. One way of making the experiment is to place some of the suspected substance in a flask along with some powdered lead acetate in order to combine with and fix any sulphuretted hydrogen that may be present, a little ether is added, and the whole is well shaken up, after which a slip of paper moistened with silver nitrate solution is suspended above the ether by attaching it to the cork with which the flask is secured, and the flask if then placed in a dark place to avoid the actinic effects of light. In from a few minutes to an hour the paper will be blackened, and will acquire a lustre from deposition of metallic silver.

Although in a dead body it is advisable to make the examination for phosphorus as early as possible, positive results may be obtained at long intervals after death. In a case that had been poisoned with phosphorus, **Hofmann**<sup>1</sup> found the head of a match in the intestines five months after death, and, with **Mitscherlich's** test, obtained satisfactory evidence from it of the presence of phosphorus. **Felletár**,<sup>2</sup> by both **Mitscherlich's** and **Dussart-Blondlot's** methods, demonstrated the presence of phosphorus in an exhumed body twelve months after death, and in another case thirteen months after death. When dealing with exhumed bodies an objection may be raised that sufficient phosphorus might be yielded by the putrefying tissues to afford indications of its presence with the above described tests, from experiments that have been made this has been found to be impossible, and, therefore, when the reactions are obtained they must be due to phosphorus derived from an extraneous source.

## IODINE.

The colour and pungent odour of iodine prevent its being used for criminal poisoning, and its exceptional presence in the household in any but small quantities—usually in the form of tincture—account for the rarity with which it is resorted to as a means of committing suicide. It is strongly irritant, and if swallowed in the solid form may produce corrosion.

**Symptoms** produced by swallowing large doses of the tincture.—Burning pain in the mouth and throat, and shortly after in the stomach, followed by salivation, vomiting, and diarrhoea. The vomited matters afford evidence of the presence of iodine, if the stomach contained starchy food at the time

<sup>1</sup> *Lehrbuch der ger. Med.*, 1887

<sup>2</sup> *Gyógyiszat*, Budapest, 1889.

the poison was swallowed, the vomit will be blue, if not, or if the iodine is in excess, the colour will be yellowish or brown. Blood has been found in both the vomit and the motions. The lips, and possibly the corners of the mouth and the chin, may be stained yellow, the mucous membrane of the tongue and mouth being whitish, the pulse is small, the surface cold and the usual symptoms of collapse are present.

Injection of strong solutions of iodine into the cavities of the body for therapeutic purposes has produced the more essential of the above-named symptoms—vomiting, with the presence of iodine in the rejected matter, thin pulse, cold, pallid surface, salivation, and dyspnoea from tumefaction of the mucous membrane of the larynx. All the mucous surfaces and the eyelids are swollen, and the skin is not unfrequently covered with an eruption. There is a tendency to heart-paralysis, which sometimes occurs a day or more after the acute symptoms have subsided. Formerly many deaths occurred from the treatment of ovarian tumours, chronic abscesses, and empyema by the injection of solutions of iodine, the proceeding is not without considerable risk when the iodine is exposed to a large absorptive surface.

Iodine is freely eliminated by the kidneys. Huber<sup>1</sup> found 0.278 gramme of iodine in 300 c.c. of urine, obtained from a woman who swallowed about four grammes of the tincture. It is also eliminated in the saliva, the milk, and in the secretions of mucous membranes.

**Fatal Dose.**—Not precisely known, as the tincture which has usually been taken is of no definite strength. One drachm of the tincture has caused death, and recovery has taken place after one ounce, calculated to contain half a drachm of solid iodine. Only eight or nine fatal cases are recorded. Death has occurred in twenty-four hours.

**Iodoform.**—The use of iodoform in dressing wounds, and its injection in solution into chronic abscesses, have caused death, grave symptoms followed by recovery not unfrequently occur. A variety of symptoms have been observed—as elevation of temperature, rapid pulse, gastro-intestinal irritation, skin-eruptions, cerebral disturbance with delirium or coma. Iodoform is most likely to produce dangerous effects when injected in the form of ethereal solution. Gaillard<sup>2</sup> records a case in which the injection into an abscess of about eighty grains, dissolved in ether, produced cessation of breathing and apparent death, artificial respiration was resorted to, and the patient recovered. Barois<sup>3</sup> records a case in which a patient died in a comatose condition on the ninth day after the injection of an ethereal solution containing forty-five grains of iodoform. Death has resulted from the free use of iodoform to dress open wounds, such as those produced by amputation of the breast or of the leg. Czerny<sup>4</sup> relates the case of a woman, aged fifty-eight, in which the wound produced by removal of a breast and the axillary glands was dressed with a drachm and a half of iodoform, three days later symptoms resembling those of meningitis set in, decubitus followed, and death occurred on the twenty-third day. Delirium, fever, and erythematous rashes have on several occasions been caused by dressing large wounds with iodoform gauze, amblyopia has been produced in the same way, and also by the internal administration of iodoform. Scotomata in the red field, the increased perception of green have been observed, with subsequent optic atrophy.

**Potassium Iodide**, when administered medicinally, occasionally produces a number of toxic symptoms, which are of therapeutical rather than toxicological

<sup>1</sup> *Zeitschr. f. klin. Med.*, 1888.

<sup>2</sup> *Bull. de Chirurg.*, 1889.

<sup>3</sup> *Arch. de Med. et de Pharm. Milit.*, 1890.

<sup>4</sup> *Wiener med. Wochenschr.*, 1882.

interest, iodism, with its accompanying skin eruptions and glandular affections, is a well-recognised condition, but as it results from medicinal treatment, and is rarely fatal, medico-legal investigation is not required. Death is stated to have occurred in one or two instances from the use of potassium iodide. Wolfe<sup>1</sup> records the case of a woman who took four six-grain doses, at intervals of four hours, which produced swelling of the face and a pemphigoid eruption, involving the mucous membrane of the nose, mouth, throat, and larynx, on the fourth day diarrhœa with blood-stained stools occurred, the woman dying on the eighth day. Conchon<sup>2</sup> records the case of a man, aged fifty-five, who, on account of an enormously enlarged thyroid, took one drachm of potassium iodide daily for fifteen days. He vomited and had diarrhœa, the action of the heart was exceedingly rapid and irregular, the pulse was very small and could not be counted. The iodide was stopped, but the symptoms increased, the goitre disappeared entirely and, although his appetite was exaggerated, the patient emaciated rapidly and died a month later. Death was probably due to action of the iodide on the thyroid, rather than to any specifically poisonous effect of the drug, it has been observed that severe poisoning by potassium iodide is most frequent in patients with goitre.

**Treatment.**—Acute poisoning by free iodine demands evacuation of the stomach with tube, or emetic, followed by farinaceous mixtures, as starch, arrowroot, flour, and the like, which have been cooked so as to rupture the starch-granules. Morphine and stimulants may be required.

**Post-mortem Appearances.**—Not well known. Yellow staining and softening of the mucous membrane of the mouth, œsophagus, and stomach have been observed, a kind of exudation-product, resembling false membrane, has been found in the larynx. Gastritis may be present, and the inflammation may advance as far as the duodenum.

**Chemical Analysis—Tests.**—If free iodine is present with organic matter, some of it may be extracted by shaking with carbon bisulphide, which acquires a rose, or violet-red colour, in accordance with the amount of iodine taken up. If the iodine is in simple combination, it may be liberated by nitric acid, and then extracted as above. If in combination in organic admixture, potassium hydroxide should be added and, after desiccation, the organic matter should be destroyed by heat, when cold the iodide is dissolved out in alcohol, evaporated to dryness, and treated with sulphuric acid, so as to set free the iodine, which is recognised by its reaction with starch.

## BROMINE.

A few fatal cases of poisoning by bromine in the liquid state are recorded. In one reported by Snell,<sup>3</sup> a man swallowed one ounce of bromine on an empty stomach, half an hour after he was found suffering from intense burning pain and eructations, there was neither vomiting, purging, nor thirst, but the patient experienced a frequent desire to evacuate the bowels. In two and a half hours symptoms of collapse occurred, and death took place seven and a half hours after the poison was swallowed. On section, the mucous membrane of the œsophagus was inflamed, the external surface of the stomach was much injected, and displayed several ecchymosed spots, internally its surface appeared like tanned leather, hard and black in colour, and could easily be peeled off, the duodenum presented the same appearance, but the mucous membrane

<sup>1</sup> *Berliner klin. Wochenschr.*, 1886.

<sup>2</sup> *Bull. et Mem. de la Soc. de Therap.*, 1895.

<sup>3</sup> *New York Journ. of Med.*, 1850.

between the valvulæ conniventes was softened, the peritoneum and omentum were stained a reddish-yellow. In a case reported by Schmalfuss,<sup>1</sup> the dead body of a man was found, the lips and tongue being dry, hard, and dark-brown in colour. On opening the abdomen the odour of bromine was perceived, the posterior wall of the stomach was altogether wanting, a portion of its anterior wall, which was grey-green in colour, being all that remained, the appearance was as though it had been burnt, a similar condition existing in the duodenum. About fifty grammes of a yellowish substance were found free in the abdominal cavity. The intestines, liver, and spleen were softened, some of the contents of the cæcum yielded bromine by simple distillation. The quantity swallowed was about ninety grammes. A third case is recorded by Herwig.<sup>2</sup> A girl, aged ten, was given by a quack a mixture containing potassium bromide, to be taken with chlorine-water. In four hours after the third dose collapse set in, followed by death in twelve hours, at the necropsy hæmorrhagic inflammation of the stomach was found. It was afterwards ascertained that each dose of the mixture, on the addition of chlorine, yielded 0.44 gramme of free bromine.

When inhaled the dense fumes given off by bromine are very irritating to the respiratory mucous membrane. Duffield<sup>3</sup> relates the case of a laboratory assistant who accidentally inhaled the fumes from about three pounds of bromine, which produced spasm of the glottis and impending death from asphyxia, by the use of steam to the throat the spasm was relaxed, and the man recovered. Kornfeld<sup>4</sup> relates a case in which a child, aged one year and three-quarters, inhaled bromine vapour, and died on the sixth day from respiratory and gastric disturbances, after death the skin of the face and neck, with which the vapour had come in contact, was parchment-like, bromine was detected in the skin and the clothing.

**Potassium Bromide.**—The evil effects of prolonged treatment with potassium bromide are not unfrequently seen, but death resulting from its use is rare. Eigner<sup>5</sup> records the case of a woman who suffered from epilepsy, for which she took potassium bromide in increasing doses until they amounted to two teaspoonfuls daily, which were continued for several weeks. She became salivated, with fætor of the breath, and inflammation of the gums, delirium supervened, and she died in five days. Dougall<sup>6</sup> saw a man, aged forty-two, who took an ounce of potassium bromide one night and half an ounce the following night. When admitted into hospital he was semi-comatose, the pulse was weak (60), and the respirations were deep, slow, and easy, without stertor, the temperature was 96.8° F, the extremities were cold and blue, and the face was livid. The pupils were of normal size. The reflexes were abolished. The patient remained two weeks without much improvement, but eventually recovered.

**Treatment.**—In the exceptional cases in which bromine is swallowed, treatment would probably be of little avail, after evacuation of the contents of the stomach, albumin or starch might be given. Poisoning with the vapour is best treated with steam inhalations.

**Chemical Analysis.**—Uncombined bromine may be separated from organic admixture by distillation, if it is in combination, it may be set free by saturating the solution with potassium dichromate, and acidulating with dilute sulphuric acid before distilling. Solid masses of organic matter may be pulped, mixed with a saturated solution of potassium hydroxide, evaporated to dryness, and the organic matter burnt off, the residue is treated with potassium dichromate and sulphuric acid, and distilled.

<sup>1</sup> *Vierteljahrsschr f. ger. Med.* (Supplement), 1889.

<sup>2</sup> *Zeitschr. f. Medicinalbeamte*, 1889.

<sup>3</sup> *American Journ. Pharm.*, 1867.

<sup>4</sup> *Friedreich's Blätter f. ger. Med.*, 1883.

<sup>5</sup> *Wiener med. Presse*, 1886.

<sup>6</sup> *Glasgow Med. Journ.*, 1893.

**Tests.**—Bromine may be recognised by its odour, it colours starch-paste yellow, and gives a yellowish-white precipitate with silver nitrate. A solution of bromine in water, added to a solution of phenol, gives a white precipitate of tri-bromophenol.

### FLUORINE.

**Hydrofluoric acid (HF)** in one or two instances has caused death.

King<sup>1</sup> records the case of a man aged forty six who drank about half a fluid ounce of commercial hydrofluoric acid. Retching and vomiting immediately occurred, the patient being in great agony. Collapse set in and death took place in thirty five minutes. Post-mortem, the mucous membrane of the mouth was white and partially denuded of epithelium. In the œsophagus were white patches and shreds of epithelium. The stomach contained a black fluid, and in parts the mucous membrane was blackened, but it was not eroded. The mucous membrane of the trachea and bronchi was reddened. Death appeared to have been caused by closure of the glottis with shreds of mucous membrane. A man aged fifty-one drank about a tablespoonful of hydrofluoric acid diluted with water. He became collapsed and died in about an hour. Post-mortem, the lips, mouth, and tongue were more or less charred and the fauces were of a deep red colour and ecchymosed. The œsophagus was slate coloured with deep-red patches. The mucous membrane of the stomach was ecchymosed, but it was not denuded, nor was there any perforation. The blood was dark and tarry. The lungs were congested and almost black in colour. Stevenson<sup>2</sup> examined the acid and found that it contained 9.2 per cent of HF, which is only one fourth the strength of the acid used by glass etchers. Schwyzer<sup>3</sup> mentions the case of a man who suffered from chronic fluorine poisoning caused by contaminated beer, and describes the result he obtained by the administration of small doses of sodium fluoride, over long periods, to animals. In both the man and the lower animals there was an increase in the myelocytes at the expense of the other leucocytes, pain in the bones, change in colour of the bone marrow from yellow to red, abnormal coagulability of the blood, diminution of the chlorides, enormous excess of calcium in the urine and faeces, lessening of the specific gravity of the bones.

The inhalation of the vapour of hydrofluoric acid, and of hydrofluosilicic acid may cause death. Cameron<sup>4</sup> records two cases in which death was thus produced. A healthy man, who worked in an artificial manure manufactory, went into a chamber which contained a quantity of freshly prepared superphosphate of lime. He became unwell, had great difficulty in breathing, and died the same evening. The following year another man, employed in the same works, was suddenly attacked with hurried and difficult breathing, and died asphyxiated in a few hours. Post-mortem, the blood was dark coloured. The lungs were œdematous, the pulmonary veins were congested and the bronchi were plugged with frothy mucus. The right side of the heart was distended and filled with clotted blood. Cameron found fluorine and silicon in the lungs, due to the inhalation of fluosilicic acid derived from the phosphates, which contained over 6 per cent of calcium fluoride.

### CHLORINE.

Fatal poisoning by chlorine is rare, the opportunities for its occurrence being limited to chemical works and bleach works. In the latter a form of chronic poisoning is met with, in which the patient acquires an anæmic or chlorotic look, loses flesh, and suffers from dyspeptic troubles associated with gastric catarrh, the sense of smell is blunted, and the bronchial mucous membrane may be affected.

The following fatal case of acute chlorine poisoning is recorded by Sury-Bienz<sup>5</sup>. A man, aged forty-eight, who worked in a chemical manufactory, accidentally took one or two breaths of pure chlorine, he at once experienced irritating cough, dyspnœa, and stabbing pain in the breast. On the following day the cough persisted and the dyspnœa was urgent, but there was but very little expectoration, the respirations were accelerated to 48, and the pulse

<sup>1</sup> *Trans. Path. Soc. Lond.*, 1873.

<sup>2</sup> *Brit. Med. Journ.*, 1899.

<sup>3</sup> *Journ. of Med. Research*, 1903.

<sup>4</sup> *Dublin Journ. Med. Sc.*, 1887.

<sup>5</sup> *Vierteljahrsschr. f. ger. Med.*, 1888.

retarded to 48 per minute. There was no albumin in the urine. The breathing became worse, the expirations being very short, and the patient died in less than forty-eight hours. On section the lungs were found to be emphysematous and œdematous, they were not consolidated, but the air-passages contained a reddish, frothy fluid, the epiglottis was pale and free from tumefaction, as was also the mucous membrane of the larynx, that of the trachea and bronchi was diffusely reddened, no fatty changes observable in the heart and other organs. Death appeared to be due to heart-paralysis, which is in accord with the results of physiological experiments. Cameron<sup>1</sup> relates the case of a man who was found dead in the fore-castle of a vessel in which chlorine had accumulated from some casks containing chlorinated lime, the appearances were those of death from asphyxia, the odour of chlorine was perceived in the ventricles of the brain.

**Bleaching Fluid**, which consists of a solution of potassium or sodium hypochlorite, with free chlorine, has been used for suicidal and homicidal purposes. The symptoms and post-mortem appearances are more marked in the digestive track than in the case in poisoning with gaseous chlorine, gastro-enteritis is produced along with the respiratory symptoms just described. Between three and four drachms caused the death of an infant, recovery has followed twenty ounces.

**Treatment.**—The dyspnoea caused by the inhalation of chlorine is best relieved by steam inhalations. Dilute sulphuretted hydrogen has been recommended on the ground that chlorine combines with the hydrogen, causing separation of the sulphur, but it is of doubtful value, the only benefit that could accrue would be the removal of any free chlorine in the air passages—which is accomplished by fresh air, sulphuretted hydrogen could not undo the damage sustained by the mucous membrane, though it might increase it. Poisoning with bleaching fluid should be treated by evacuation of the stomach, by demulcents, and morphine.

## BORON.

**Boracic Acid** [B(OH)<sub>3</sub>], or boric acid, is used surgically as an antiseptic, and commercially as a preservative of milk and other articles of food, it has very little taste, hence its use as a food preservative. When experimentally administered to animals, boracic acid produces prostration, feebleness of the pulse, and diminution of the respiratory activity, parenchymatous nephritis, cloudy swelling with fatty degeneration of the epithelium, and hæmorrhages under the capsule of the kidney have been observed.

Fatal poisoning has followed the injection of solutions of boracic acid into natural and into abscess-cavities of the body. Molodenkow<sup>2</sup> relates two such cases. In one a 5 per cent solution was injected into the pleural sac on account of an empyema, vomiting occurred and the pulse was small and weak, the following day erythema appeared on the face and spread over the body, on the third day the patient died. The second case was that of a boy who half an hour after having an abscess-cavity washed out with a boracic solution began to vomit and to be collapsed, erythema and hiccup occurred on the second day, when he died. The necropsy yielded negative results, except that there were a few ecchymoses on the inner surface of the pericardium, death was due to heart-paralysis. Three fatal cases from washing out the stomach

<sup>1</sup> *Dublin Quarterly Journ. of Med. Sc.*, 1870

<sup>2</sup> *Petersb. med. Wochenschr.*, 1881.

with a solution of boric acid are recorded by Hogner<sup>1</sup> The symptoms produced were —General depression, erysipelatous eruption on the face and purpuric spots on the body, elevated temperature, vomiting, diarrhoea, frequent desire to urinate, blood in the urine, stupor, and death in one case on the third day Welch<sup>2</sup> records a case of poisoning from the use of a vaginal tampon of boric acid —Formication of hands and feet, swelling of the skin of the face, hands, and feet, with pronounced depression of the nervous system, dysuria, prostration and collapse ensued Recovery took place, the skin undergoing general desquamation Best<sup>3</sup> records a fatal case which was due to tamponing a wound in the inguinal region with boric acid The symptoms produced comprised skin rash, cyanosis, collapse, rapid, irregular pulse, rapid respiration, excessive vomiting and elevation of temperature to 100.8° F The affection of the skin is one of the most constant signs of boric poisoning Lemoine<sup>4</sup> reports four non-fatal cases from surgical practice —Erythema and urticaria were present in all, among the other symptoms were vomiting, delirium, hallucinations, and in one case diplopia Wild<sup>5</sup> records several cases of dermatitis caused by the internal use of boric acid and borax Sanders<sup>6</sup> describes a case following rectal injections of boric acid Noisy delirium and a rash consisting of hard shotty papules, subsequently becoming purpuric, developed

As a food preservative boric acid is usually employed in combination, in the form of borax Although it may be difficult to trace any ill effects to the swallowing of small doses of borax, there are good grounds for assuming its noxiousness, especially when added to milk which constitutes the chief food of young children

The elimination of boric acid takes place chiefly through the kidneys Rost<sup>7</sup> shows that after reception by the mouth, 50 per cent is excreted in the urine within the first twelve hours, the remaining 50 per cent takes six or eight times as long The maximum output occurs two to three hours after reception, traces may often be obtained as long as nine days after Rost gives a copious and most useful "literature" on boric acid The continued reception of small amounts of borax with food tends to cause diarrhoea, the assimilation of the food is diminished, and emaciation or loss of weight results

**Chemical Analysis.**—Organic fluids containing boric acid, or borax, may be evaporated down, treated with sulphuric acid, and extracted with alcohol

**Tests.**—The alcoholic extract of boric acid burns with a green-coloured flame Boric acid partially reddens blue litmus paper and browns tumeric paper, the reaction with tumeric is distinguished from that due to alkalis by not disappearing under the influence of acids

<sup>1</sup> *Eva*, 1884.

<sup>2</sup> *New York Med Rec*, 1888

<sup>3</sup> *Trans Chicago Path Soc*, 1905

<sup>4</sup> *Gaz Med. de Paris*, 1890

<sup>5</sup> *The Lancet*, 1899

<sup>6</sup> *Brit Med Journ*, 1912

<sup>7</sup> *Arch intern de Pharmacodyn*, 1905

## CHAPTER XXXII

## GASEOUS COMPOUNDS.

## SULPHURETTED HYDROGEN.

POISONING with pure sulphuretted hydrogen ( $H_2S$ ), excepting in chemical works, is rare, in those cases in which mischief results from its inhalation, the mixed gas known as **sewer gas** is the combination in which it usually exists. Sewer gas is composed of a variable mixture of sulphuretted hydrogen with free hydrogen, carburetted hydrogen, ammonia, carbon dioxide, and atmospheric air deprived of part of its oxygen. Although several of these gases are poisonous, the toxic effects of sewer gas are chiefly due to the sulphuretted hydrogen it contains, one description of the symptoms and post-mortem appearances, therefore, will serve both for sulphuretted hydrogen and for sewer gas.

Much of the  $H_2S$  in sewer gas is probably produced by the action of bacteria on the albuminous substances present in the sewage, this only occurs, however, when the air in the sewer is stagnant, or in other words, in the absence of oxygen, if the sewer is efficiently ventilated, the bacteria oxidise to sulphates the sulphur which is split off, and little or no  $H_2S$  is formed. Hoppe Seyler<sup>1</sup> found that, when solutions containing albumin undergo decomposition in the presence of free oxygen, the only volatile products are carbon dioxide, ammonia, and water. Lehmann's<sup>2</sup> experiments show that an atmosphere containing 0.05 per cent of  $H_2S$  produces in man alarming symptoms in a few minutes, 0.2 per cent is rapidly fatal to cats and dogs. It is to be borne in mind as regards  $H_2S$  that the lowering of a lighted candle into a sewer, or manhole, yields no information as to the respirable condition of the contained air, the light will go out if the oxygen is reduced by 3 or 4 per cent., and yet such an atmosphere may be breathed without inconvenience. On the other hand, the candle will continue to burn in an atmosphere containing a percentage of  $H_2S$  that would prove rapidly fatal to human beings. Air stagnant over sewage, containing much  $H_2S$  in solution may become poisonously charged with the gas, whilst the air of a well-ventilated sewer containing like sewage will be innocuous. Haldane<sup>3</sup> analysed some air taken from a sewer three days after the occurrence in it of fatal gaseous poisoning, and merely found a diminution of 0.07 per cent of oxygen with an increase to the same amount of carbon dioxide, yet some of the sewage, taken at the same time from this sewer, rendered poisonous air which was allowed to stand over it for a short time. Fatal accidents, due to rapid liberation of  $H_2S$ , have happened owing to workmen stirring the sewage contained in a manhole or well, in one instance the discharge of a quantity of dilute sulphuric acid into a sewer suddenly liberated a poisonous amount of  $H_2S$  from the sulphides which were present.

**Symptoms.**—When only a limited percentage of the gas is present it nevertheless acts as an irritant to the mucous membrane of the air-passages, and causes a sensation of "catching of the breath", the respirations are rendered slow and difficult, the pulse becomes small, there is a feeling of oppression in the head, accompanied by sickness, dizziness, and probably diarrhoea, at the same time great muscular prostration is experienced. If the percentage is large, urgent symptoms of asphyxia and heart failure, with profound collapse, cyanosis, dilated pupils, unconsciousness, delirium, and convulsions may occur.

<sup>1</sup> *Zeitschr. f. physiol. Chemie*, 1884.<sup>2</sup> *Arch. f. Hygiene*, 1892.<sup>3</sup> *The Lancet*, 1896.



Beyond conjunctivitis and broncho-pneumonia, secondary effects are very rare. Wilesworth<sup>1</sup> records the case of a man employed in a chemical works who became maniacal, and continued so for two or three weeks, after having accidentally inhaled sulphuretted hydrogen, he began to improve at the end of a month, but did not recover his mental vigour for five months after his admission into the asylum, which took place about a week after the onset of the symptoms.

Asphyxia, due to the action of the gas on the hæmoglobin, and probably also on the tissues, by which they are respectively rendered incapable of yielding up and of receiving oxygen, is believed by some to be the cause of death, others attribute death to certain disturbances of the nervous system by which the pulmonary and the cardiac innervation is deranged. Kaufmann and Rosenthal<sup>2</sup> demonstrated experimentally that inhalation of sulphuretted hydrogen lowers the blood-pressure, and lessens cardiac action by stimulation of the vagus centre. Brouardel and Loye<sup>3</sup> found that in animals which are made to respire sulphuretted hydrogen the pupils are dilated, the heart-beats slowed, and the respirations are gradually diminished in amplitude, in some cases the heart continued beating for two minutes after respiration had ceased. Pohl<sup>4</sup> believes that the presence of sulphuretted hydrogen in the blood determines the formation of sodium sulphide which causes paralysis of the central nervous system. Lehmann<sup>5</sup> thinks that in animals death is not solely due to changes in the blood and to paralysis of the central nervous system, but also to œdema of the lungs. Ushinsky<sup>6</sup> infers from experiments on animals that the toxic action of  $H_2S$  cannot depend on the formation of sulphur-methæmoglobin, since large quantities of blood charged with the gas can be injected into the circulation without causing the least harm, although sulphur-methæmoglobin can readily be distinguished in blood subsequently withdrawn from the animal, and further, that in animals which have been poisoned with  $H_2S$  sulphur-methæmoglobin cannot always be detected, he considers death to be solely due to paralysis of the central nervous system.

It was formerly taught that  $H_2S$  is freely eliminated by the lungs, but more recent investigations are against this view, Laborde<sup>7</sup> found that a residuum of the gas remains in the blood after it has passed through the lungs, Ushinsky states that  $H_2S$  is only feebly eliminated by the lungs.

**Treatment.**—In order to combat the tendency to death, and at the same time promote elimination, artificial respiration should be vigorously carried out. Cold effusion has been recommended, but if the surface is already cold it would be worse than useless, the application of external warmth would rather be indicated. Cautious inhalation of chlorine diluted with air has been suggested on the ground that it combines with the hydrogen of the  $H_2S$  and precipitates the sulphur, it might be very carefully tried along with artificial respiration.

**Post-mortem Appearances.**—Putrefactive changes quickly follow death. In some cases cadaveric rigidity is reported to have been well marked, which is contrary to what might be expected, as after poisoning by sulphuretted hydrogen the molecular vitality of the muscles disappears with the occurrence of somatic death. The blood is fluid and dark in colour, and in consequence the organs which are rich in blood—as the liver, lungs, and spleen—are also

<sup>1</sup> *Brit Med Journ*, 1892

<sup>2</sup> *Arch f Anat u Physiol*, 1865

<sup>3</sup> *La France Médicale*, 1885

<sup>4</sup> *Arch f exper Path*, 1887

<sup>5</sup> *Arch f Hygiene*, 1892

<sup>6</sup> *Zeitschr f physiol. Chemie*, 1892.

<sup>7</sup> *Comptes rendus de la Société de Biologie*, 1886

darker than usual. The brain has been found of a peculiar dirty greyish-green owing to the colour of the blood, the muscles are dark, including those of the heart, they sometimes present a bluish tint. The lungs will probably be œdematous. The remaining appearances are common to other forms of death from asphyxia.

Sulph-methæmoglobin shows a narrow band towards the red end of the spectrum, between C and D, resembling that of methæmoglobin, but it does not disappear on the addition of a reducing agent as the methæmoglobin band does when similarly treated. The blood of human beings poisoned by  $H_2S$  has been examined spectroscopically by many observers with negative results. Rømer<sup>1</sup> saw a workman in a tar-distillery who was poisoned by  $H_2S$  whilst cleaning out a still, he was cyanosed, the pulse being 130 and the temperature  $99.5^{\circ} F$ . Blood obtained by venesection had a strong tendency to coagulate, but neither chemically nor spectroscopically could the presence of  $H_2S$  be ascertained. The patient died in about sixteen hours. The autopsy revealed advanced putrefaction, the organs were deep red in colour and there was diffuse fatty degeneration of the heart with œdema of the lungs. Laborde, Uschinsky, and others, in experimental poisoning of animals with sulphuretted hydrogen, obtained (but by no means invariably) the characteristic spectrum of methæmoglobin in combination with sulphur. Uschinsky and Binet<sup>2</sup> found that the blood of frogs so poisoned nearly always yields the spectrum of sulph-methæmoglobin.

**Chemical Analysis- Tests.**—The odour is sufficiently distinctive to indicate the presence of even minute amounts of sulphuretted hydrogen. A piece of white filtering paper dipped in a solution of lead acetate, is rapidly discoloured when suspended near tissues or other substances impregnated with the gas.

### CARBON DIOXIDE.

Poisoning with carbon dioxide ( $CO_2$ ) occurs in deep wells and excavations, in brewers' vats, and in the neighbourhood of lime-kilns and brick-kilns which are in operation, it is also one of the constituents of "after damp," the mixture of gases formed by explosives in coal mines (*q v p 442*). The cellars of houses in the neighbourhood of mines have occasionally been charged with  $CO_2$ , especially during a hard frost, by the pent-in gas passing through the porous soil beneath the foundations. Biggam<sup>3</sup> relates an instance in which seven people were poisoned in this way, of whom two died. The percentage of carbon dioxide in atmospheric air that will cause fatal results to human beings who respire it, is not accurately known, under ordinary conditions probably 20 per cent. and even much less, would soon prove fatal. Human beings can breathe air mixed with 20 per cent. of *pure*  $CO_2$  for some time without life being endangered, but a much lower percentage of the gas as it is exhaled from the lungs is lethal, within limits, both men and animals may acquire a certain tolerance for the gas, and breathe air contaminated with it that would be noxious to the untrained organism. The test usually applied to ascertain whether air charged with  $CO_2$  is respirable or not is to lower a lighted candle into the mixed gases, if it goes out the atmosphere is poisonous. To this extent the test is trustworthy, but the converse is not to be assumed—namely, that if the candle burns the air is harmless, a candle will burn in a percentage of  $CO_2$  that is dangerous to life.

<sup>1</sup> *Munchener med Wochenschr*, 1897.

<sup>2</sup> *Revue Med. de la Suisse rom.*, 1896.

<sup>3</sup> *Brit Med Journ.*, 1893.

**Symptoms.**—When a poisonous but not concentrated mixture of carbon dioxide and air is respired, heaviness in the head with giddiness, noises in the ears, a sensation of tightness across the chest, and an inclination to sleep are experienced, shortly after, the muscles lose power, and consequently the individual, if standing at the time, falls to the ground. The subsequent symptoms are those of asphyxia, coma, stertorous breathing, cyanosis, and possibly convulsions, sometimes delirium occurs. A concentrated atmosphere of carbon dioxide, such as is occasionally encountered by workmen who are lowered by a rope into a well or vat which is filled with the gas, produces immediate loss of consciousness and of muscular power, death rapidly ensuing unless the victim is at once rescued.

When carbon dioxide diluted with air is respired, it acts both as a poison and also—when the partial pressure of the gas in the atmosphere is greater than that in the lungs—as a preventive to excretion of the physiologically formed  $\text{CO}_2$ . The ultimate effects on the organism, allowing for the degree of concentration in which  $\text{CO}_2$  is respired, are the same as those of asphyxia resulting from the cutting off of oxygen and the consequent accumulation of carbon dioxide in the tissues, but the respiratory movements more quickly cease than when the respired air is simply deficient in oxygen, in the latter case the excretion of  $\text{CO}_2$  is but little affected.

The **treatment** is that for asphyxia—artificial respiration, external warmth and stimulants.

**Post-mortem Appearances.**—They are simply those of death from asphyxia, dark-coloured, fluid blood, fulness of the right heart and veins, usually with hyperæmia of the lungs, and frothy mucus in the air-passages.

**Chemical Analysis.**—The analysis that may be required is that of the atmosphere in which the poisoning took place. A sample may be obtained by displacing the air in a dry flask or bottle of five or more litres capacity when immersed in the suspected atmosphere, this may be done with a pair of bellows, or if the gas is contained in a well or other excavation the flask, filled with fine dry sand, is attached to a cord and let down to the required depth, by means of a second cord fastened to the lower part of the flask it is inverted, so that the sand flows out and is replaced by gas, the flask is withdrawn mouth upwards and immediately stoppered. The amount of  $\text{CO}_2$  in the flask is ascertained by adding 20 to 50 c.c. of a titrated solution of barium hydroxide, replacing the stopper and shaking well for some minutes, the loss of hydroxide by conversion into carbonate, which falls as a white precipitate, is estimated by titrating the solution with oxalic acid.

## CARBON MONOXIDE.

In medico-legal practice carbon monoxide (CO) is met with in the fumes given off by slow combustion stoves, or fire-places, in which the exit of the gaseous products of combustion is restricted, in admixtures of atmospheric air with coal-gas or water-gas and mixed with nitrogen and carbon monoxide in the gases formed by the explosives used in mines, death has resulted from sleeping in a closed room, in the open fire-place of which coke instead of coal was burnt. Poisoning by CO not unfrequently causes the death of those who are unable to escape from a building which is on fire.

In all these instances other gases besides CO and air are present, but the toxic effects are chiefly, if not entirely, due to the CO. It is impossible to determine the minimum percentage of CO in air that will cause death to human beings, according to Haldane,<sup>1</sup> 0.05 per cent in otherwise normal air produces distinct toxic symptoms, and with about 0.2 per cent urgent symptoms are

<sup>1</sup> *The Journal of Physiology*, 1995.

produced, 1 per cent is usually accepted as a fatal admixture. Coal-gas contains a variable amount, from 4 to 7 or 8 per cent of CO, which is practically the sole poisonous component, along with varying amounts of hydrogen, carburetted hydrogen, watery vapour, nitrogen, and carbon dioxide. Coal-gas is easily recognised by its peculiar odour, which betrays its presence in the air in amounts of only 0.01 to 0.02 per cent, it has been observed, however, that when coal-gas is allowed slowly to percolate through a thick layer of earth (6 or 7 feet) it may become almost odourless. Biefel and Poleck<sup>1</sup> show, by analysis of coal-gas after passing through earth, that the loss of odour is due to absorption of the greater part of the heavy carburetted hydrogen and marsh gases, and further, that the percentage of CO is increased. It occasionally happens that the inmates of a house in which the gas fittings are perfect or even in one which is not supplied with gas, are poisoned by coal-gas which has percolated through the ground from a fractured street-main which may be at some distance, in one such instance the seat of fracture was 86 yards away from the house into which the gas escaped. It is to be noted that a mixture of coal-gas with air is poisonous before it reaches the explosive point, Jones<sup>2</sup> saw two persons who were poisoned (one fatally) by the escape of coal-gas into a room on the dressing-table of which a paraffin lamp was found burning. Water-gas, which is sometimes used as a substitute for, or an adjunct to, coal-gas, contains as much as 10 per cent of CO, its dangerous properties being accentuated by the absence of the characteristic odour. Possessing feeble illuminating value, water-gas requires enriching with some volatile or gaseous hydrocarbon to fit it for use as a domestic illuminant, the favourite way is to mix it with oil-gas, produced by allowing mineral or other oil to pass into heated retorts in which it is decomposed into a more or less permanent gas rich in hydrocarbons. The so-called water-oil-gas, or carburetted water-gas, contains 20 per cent of CO, when mixed with coal-gas the combined gases contain from 14 to 16 per cent of CO, in either case the consumers are supplied with an unnecessarily poisonous illuminant.

The use of gas burners on the Bunsen principle to heat large bodies of cold water, as is done in some bath-rooms, is attended with considerable risk, the flame is rapidly cooled by contact with the surface of the water reservoir, and combustion is consequently rendered imperfect, the oxides of carbon—especially the monoxide—with acetylene being given off, when water is heated in this way, provision for abundant ventilation should be made. Gasoline stoves, if used in small rooms, are not free from danger. M'Cormick<sup>3</sup> relates a case in which a man and his wife were found dead in a bedroom heated by a gasoline stove, the gas given off being principally carbon monoxide.

In one respect carbon monoxide constitutes an exception to the poisons which naturally exist in the gaseous form, on the Continent, especially in France, it is frequently used for suicidal purposes, a mode of self-destruction that is almost unknown in England. The usual way is for the suicide to shut himself up in a closed room along with a receptacle containing ignited coke, or charcoal, a method which is often adopted by two persons who desire to die simultaneously. Coal-gas is often used as a suicidal agent, a man took to bed with him an unlighted Bunsen burner which was connected with the main by means of an elastic tube, he inhaled the gas, and when he was discovered death had taken place. In America, where water-gas is freely used for illuminating purposes, its use as a suicidal agent is not uncommon.

Carbon monoxide poisoning occurs in two forms—**acute** and **chronic**.

<sup>1</sup> *Zetschr f Biologie*, 1880

<sup>2</sup> *Brit Med Journ*, 1896

<sup>3</sup> *Med News Phil*, 1891

**Acute Poisoning by Carbon Monoxide.**

**Symptoms.**—There may be a preliminary period of excitation, which is quickly followed by a sensation of heaviness in the head, dizziness, noises in the ears, accelerated cardiac and respiratory movements, oppression on the chest, and occasionally nausea and vomiting. Along with these symptoms muscular weakness occurs, with drowsiness, loss of sensation and of the reflexes, to which coma succeeds, in fatal cases convulsions frequently precede death. The pulse is small and becomes more so as the gravity of the case increases, so that when the patient is first discovered, the radial pulse is often imperceptible. In the state of coma the conjunctivæ are intensely hyperæmic, and the eyes have a staring appearance, with partially dilated and insensible pupils. The whole of the skeletal muscles, including the sphincters, are relaxed, the surface, especially of the limbs, is cold and cyanotic, and the lips are often covered with froth. In less severe cases slight elevation of temperature has been observed. On rare occasions exceptional symptoms have occurred, Caspar states that a man was suddenly attacked with transitory mania after inhaling air contaminated with CO.

Shufflebotham<sup>1</sup> has described the symptoms which were presented by many of the men who were rescued from the Senghenydd colliery after a serious explosion. In those who had been badly "gassed" the pulse-rate was most irregular, not only as regards frequency but as regards volume and character, and examination of the pulse gave no indication of the physical condition of the patient. At one time it might be slow, feeble and irregular, apparently the pulse of a person slowly sinking, but without apparent cause it would ten minutes later be of normal rate and materially improved in quality. The majority of cases showed large erythematous patches of a bright cherry colour with clearly defined margins but of irregular shape. They were tender and tense with a considerable amount of induration, and were at first regarded as burns. Six weeks after the explosion the patches were smaller and the induration had subsided to a large extent, but not entirely. The skin over the patches was discoloured and there was desquamation, while the follicles were prominent. The œdema around the patches had disappeared and there was hyperæsthesia over the patch of skin which remained and anæsthesia over the area of skin which had been originally affected. In many cases there was peroneal palsy, or ankle drop, developing from two to four days after the explosion. The earlier symptoms were pains in the legs with tenderness along the peroneal nerve and talipes equino-varus. In some cases it appeared as though the paralysis would be permanent. Six weeks after the explosion the symptoms were less of power and dorsiflexion of the foot. In one or two instances there was complete paralysis in this respect, and in some diminished power of plantar flexion. The cutaneous reflexes were absent over the area of skin supplied by the peroneal nerve, and in some cases irregular areas of skin apart from the distribution of this nerve were similarly affected. Pneumonia occurred in several cases. Retention of urine was a striking symptom in every case, and in two cases there was paralysis of the sphincter ani.

It has been frequently stated that sugar is almost invariably present in the urine of those suffering from poisoning by carbon monoxide, its occurrence, however, is only occasional. Maschka<sup>2</sup> in two out of twelve cases found a trace only of sugar in the urine. Hoppe-Seyler<sup>3</sup> constantly found a substance

<sup>1</sup> Milroy lectures, 1914<sup>2</sup> *Prager med. Wochenschr.*, 1880<sup>3</sup> *Physiolog. Chemie*, 1881

in the urine which reduced copper salts, but never any glucose Garofalo<sup>1</sup> examined the urine from a number of dogs poisoned with carbon monoxide, and failed to find a trace of sugar

In medico-legal practice the symptoms observed in poisoning by carbon monoxide are not always exactly the same, this arises from the admixture of other gases Poisoning by pure CO can only be procured experimentally, but the salient features of CO poisoning are present in all cases in which that gas is the principal toxic component

Carbon monoxide is a typical example of a cumulative poison, its powerful toxic action being due to its affinity for hæmoglobin, which is nearly 140 times, or according to Dresser,<sup>2</sup> 200 times, greater than the affinity of oxygen for hæmoglobin When CO is inhaled it gradually displaces the oxygen of the hæmoglobin, with which it combines, forming carbon monoxide hæmoglobin, or carboxyhæmoglobin—a much more stable compound than oxyhæmoglobin, according to Hufner,<sup>3</sup> the dissociation-constant of carboxyhæmoglobin is about 33 times less than that of oxyhæmoglobin under like conditions The combination is sufficiently intimate to resist the action of reducing agents, but it gradually yields to the action of oxygen, if air or oxygen is passed for a long time through a solution of carboxyhæmoglobin the CO is gradually separated from the hæmoglobin, O<sub>2</sub> taking its place In the living body carbon monoxide hæmoglobin neither takes up nor gives off oxygen, it is, therefore, incapable of acting as an oxygen carrier to the tissues, death usually occurring before the whole of the hæmoglobin is saturated with CO Upon the degree of saturation depends the possibility of recovery, if a sufficient amount of hæmoglobin remains free as to enable internal respiration to be carried on in such a way as to maintain life until the combined CO is gradually dissociated, recovery is possible, if not, death takes place from asphyxia When, as is usually the case, a limited percentage of CO is present in the air, the symptoms do not manifest themselves until after the individual has breathed the contaminated air for some time According to experiments made by Haldane,<sup>4</sup> it is necessary, in human beings, that the blood should be one-third saturated before the characteristic symptoms develop when the blood is half-saturated the symptoms become urgent This explains the delay in the appearance of the symptoms time is required to expose the blood in its passage through the lungs to the action of the gas, inasmuch as each inspiration of contaminated air contains only a small percentage of CO, and further, only about one-half of the inspired CO is absorbed Although the affinity of O<sub>2</sub> for hæmoglobin is much less than that of CO, still the presence of O<sub>2</sub> in the respired air exercises a certain inhibitory influence on the formation of carboxyhæmoglobin Dresser,<sup>5</sup> when experimenting with rabbits, found that death occurs when the capacity of the blood to take up O<sub>2</sub> is reduced 30 per cent below the normal

In carbon monoxide poisoning the blood presents a very different appearance from that which is met with when death from asphyxia results in the usual way in ordinary asphyxia the blood is dark, in CO poisoning it is bright red. This is due to carboxyhæmoglobin being irreducible, it retains its colour under circumstances that would deprive oxyhæmoglobin of its oxygen and cause it to assume the dark appearance of reduced hæmoglobin

The affinity of CO for hæmoglobin is not limited to living blood, but, the circulation having ceased, the formation of carboxyhæmoglobin in the dead body is restricted to that

<sup>1</sup> *Glicosuria per Ossido di Carbonio*, 1891

<sup>2</sup> *Arch f exp Pathol*, 1891

<sup>3</sup> *Arch v f Anat v Physiol*, 1895

<sup>4</sup> *The Journal of Physiology*, 1895

<sup>5</sup> *Arch f exp. Pathol*, 1891

portion of the blood which is accessible to the gas. An illustration of the way in which a dead body may acquire the external appearance of carbon monoxide poisoning was afforded in a case of instantaneous asphyxiation, by sulphuretted hydrogen, of two men in a sewer in Manchester. The body of one man was recovered soon after death, the body of the second man remained five days in the sewer, when it was recovered six miles from the fatal spot, having been carried along by the sewage. The first body presented the usual appearances of asphyxial death: the face was turgid and blue in colour, and the veins of the neck were gorged with dark coloured blood. The rest of the body was pallid. The second body presented, externally, the characteristic appearance of poisoning by carbon monoxide: it was rosy red all over. Internally, however, the appearances were like those yielded by the first body: the blood, muscles and viscera were dark coloured. It was evident that the rosy colour of the surface was due to percutaneous transudation of carbon monoxide after death. The condition was reproduced experimentally by luting the mouth of a glass funnel on to the surface of a cadaver and attaching the tube of the funnel to a gas supply for twelve or more hours. When the funnel was removed the underlying surface was rosy red. The presence of a small amount of carbon monoxide in the sewer was accounted for by leakage from an adjacent gas main.<sup>1</sup>

It has been stated that carbon monoxide possesses an intrinsic toxic action in addition to its power of depriving the tissues of oxygen, from experiments Linossier<sup>2</sup> deduces that CO does possess such an action, but that it is very feeble. On the other hand, Haldane<sup>3</sup> conclusively shows that CO acts solely by combining with hæmoglobin and in no other way. First, by placing animals in oxygen at a pressure of two atmospheres the blood takes up in simple solution sufficient oxygen as to render the animal independent of its red blood corpuscles as oxygen carriers, CO at a pressure of one atmosphere is then added, under these conditions the toxic action of CO is abolished although the hæmoglobin of the animal may become saturated with it. Second, by placing in an atmosphere of 75 per cent of CO and 25 per cent of oxygen animals which have no hæmoglobin, such animals are unaffected. Heineke<sup>4</sup> states that coal-gas, along with other poisons, produces ferment-intoxication in the blood, and in consequence the white corpuscles develop a tendency to adhere to each other and to form thrombi.

**Treatment.**—When a person suffering from carbon monoxide poisoning is made to respire pure air, the carboxyhæmoglobin tends to part with its CO by dissociation in the capillaries of the lungs, followed by diffusion of the gas through the alveolar epithelium into the air-passages. The process of dissociation is slow, therefore artificial respiration should be perseveringly maintained, combined, when possible, with the inhalation of oxygen. If, when diluted with about 100 drops of water, a drop of blood obtained from the finger of the patient is distinctly pinker than a similar dilution of normal blood, oxygen will be beneficial. Next in importance is the application of warmth externally. Stimulants are useful, if the patient cannot swallow they may be administered by the rectum, or ether may be injected under the skin. Two cases are recorded in which subcutaneous injections of nitro-glycerine were followed by recovery, in one, the pulse improved, and the respirations deepened forthwith. Venesection with transfusion has been tried out of twenty-three cases in eight only was the treatment attended with success. Stocker<sup>5</sup> records a very encouraging case in which transfusion was used. A man slept in a room heated by a stove, and was found next morning insensible and apparently dying. Ether injections, artificial respiration, and electrical stimulation of the phrenics were tried for forty-eight hours without effect, at least 800 grms of blood were withdrawn from the median vein, and replaced by 110 grms of defibrinated human

<sup>1</sup> *Med Chron*, 1899.<sup>2</sup> *Lyon Medical*, 1889.<sup>3</sup> *The Journal of Physiology*, 1895.<sup>4</sup> *Deutsches Arch f klin Med*, 1887 8.<sup>5</sup> *Correspondenz-Blatt f schweizer Aerzte* 1888.

blood, in two hours gradual improvement set in but the pulse, respiration, and temperature did not become normal until the third day, the ultimate recovery was prolonged over many weeks. When transfusion is resorted to, it should be preceded by withdrawal of blood, and the transfused fluid should be human blood—in other words, an appropriate oxygen-carrier—saline solutions are useless. The views of Heineke are opposed to transfusion as defibrinated blood, injected into the circulation, would tend by the introduction of a certain amount of fibrin ferment to increase the evil.

**Post-mortem Appearances.**—The external appearance is very characteristic, on account of the bright pink colour of the post-mortem stains, cadaveric rigidity is usually well marked, and passes off slowly. **Internally**, the colour of the tissues is equally striking.—The blood is cherry-red in colour, and is for the most part fluid, the blood-vessels are dilated, and, being filled with the bright-red blood, impart a very characteristic appearance to many of the viscera. Microscopically examined, the red corpuscles show no change. The brain and membranes may be hyperæmic but they often contain no excess of blood, serous effusion is not unfrequently found in the cerebral ventricles. The lungs may be hyperæmic, they have been found œdematous. The mucous membrane of the trachea and bronchi often presents a normal appearance, but it may be coated with froth. Another characteristic feature has been frequently observed to follow death from poisoning by carbon monoxide—the organs and the blood are highly resistant to putrefactive changes. Stevenson<sup>1</sup> states that portions of the liver from a case of poisoning by water-gas showed an unchanged aspect and retained the odour of the fresh organ two months after they were removed from the body, although no preservative was used, the stomach and duodenum in parts were also unchanged in appearance.

**Spectroscopic Examination of the Blood.**—When blood is fully saturated with CO (all the hæmoglobin being converted into carboxyhæmoglobin), the absorption spectrum yielded by it consists of two bands resembling those of oxyhæmoglobin, but they are slightly nearer the violet end of the spectrum, the change of position, however, is only appreciable by direct comparison of the two spectra side by side. If this constituted all the difference, it would be insufficient to afford convincing evidence for medico-legal purposes, but a further and more decided difference is manifested on the addition of a reducing agent such as ammonium sulphide—the bands of carboxyhæmoglobin are unaltered, which is in marked contrast with the change that takes place in oxyhæmoglobin when similarly treated. As previously stated, death usually occurs before the whole of the hæmoglobin has been converted into carboxyhæmoglobin, when this is the case, a mixture of carboxyhæmoglobin and oxyhæmoglobin is present in the blood, and consequently the addition of a reducing agent does not affect the hæmoglobin which is in combination with CO, but it reduces that which is in combination with oxygen. Thus the spectrum yielded by the blood of a person who has died from CO poisoning does not necessarily remain unchanged on the addition of a reducing agent, the hæmoglobin which is combined with oxygen is reduced, and shows the broad band of reduced hæmoglobin, on which are superimposed the two persistent bands of that portion of the hæmoglobin which is combined with CO (*vide* Diagram of Blood Spectra). Kunkel<sup>2</sup> states that when the amount of carboxyhæmoglobin present in the blood is below 28 per cent, the carboxyhæmoglobin bands, after the addition of a reducing agent, are not visible as two separate bands, only the broad band of reduced hæmoglobin being seen, with 30 per cent and

<sup>1</sup> *Guy's Hosp. Reps.*, 1890

<sup>2</sup> *Sitzungs- u. d. phys.-med. Gesellsch. zu Würzburg*, 1888



upwards the carboxyhæmoglobin bands appear distinctly separated. It is important to note that in an atmosphere containing a *large* percentage of CO death may occur before the hæmoglobin becomes sufficiently charged with CO as to afford spectroscopic evidence of its presence.

Hoppe Seyler's test of adding a solution of sodium hydroxide to CO blood yields a cinnamon red, normal blood is converted into a dirty brownish green mass. Salkowski<sup>1</sup> has modified this test by diluting the blood with distilled water to twenty times its volume, placing the solution in a test tube, and adding an equal volume of a solution of sodium hydroxide (S G 1.34), the solution containing carbon monoxide blood, after a momentary turbidity, becomes bright, and light red in colour, that containing ordinary blood changes to dirty brown. A very delicate test for the presence of carboxyhæmoglobin in blood is that devised by Kunkel. A little of the blood is diluted with ten volumes of water, and some 3 per cent aqueous solution of tannin is added, a precipitate is formed which is pinkish-white if carboxyhæmoglobin is present and brownish white (like *café au lait*) with normal blood. This test will detect carboxyhæmoglobin when it represents only 20 per cent of the total amount of blood, it is applicable to blood which is undergoing decomposition. Carboxyhæmoglobin is very stable. Landois obtained spectroscopic reactions from the blood of a woman in whose body general putrefaction had taken place, death from CO poisoning having occurred eighteen months previously.

**Quantitative Estimation.**—A convenient method of ascertaining the quantity of CO contained in the blood is that adopted by Grehan<sup>2</sup>. The blood to be examined is placed in a flask connected with an apparatus for the extraction of gases, some glacial acetic acid is added, and the flask is placed in boiling water, the result is that the hæmoglobin is converted into hæmatin, with liberation of the carbon monoxide. The carbon dioxide that is given off is absorbed by potassium hydroxide, and the oxygen by pyrogallol, the remainder consists of a mixture of nitrogen and carbon monoxide, the relative proportion of the two gases is arrived at by introducing a small quantity of a solution of copper chloride in hydrochloric acid, which completely absorbs the carbon monoxide.

Dresser<sup>3</sup> makes use of Hufner's spectrophotometer to determine the percentage of carboxyhæmoglobin in blood. This method is founded on the relative constancy of the absorption of light at two selected parts of the spectrum by blood from the same animal species, thus applies both to oxyhæmoglobin and to carboxyhæmoglobin. The extinction coefficients of each are calculated in wave lengths, and from certain known data previously determined by experiments, the relative proportions of oxyhæmoglobin and of carbon monoxide hæmoglobin are deduced.

Haldane<sup>4</sup> adopts a colorimetric method. He places in a small tube a 1 per cent solution of the blood to be examined, in a second similar tube is a 1 per cent solution of normal blood and in a third a like dilution of blood saturated with CO. A standard solution of carmine is then added to the normal blood until its tint resembles that of the blood which is being tested, then the amount of carmine necessary to produce equality of tint with the blood saturated with CO is ascertained, and from the data thus obtained the percentage of CO in the first tube is calculated.

### Chronic Poisoning by Carbon Monoxide.

It is probable that some of the less toxic effects of carbon monoxide are of more common occurrence than is generally suspected. Chronic CO poisoning occurs in those who work for long hours in small or imperfectly ventilated rooms heated by slow combustion stoves, or gas stoves which give off CO in too small quantity to produce immediate effects, furnacemen and stokers are subject to the risk of repeatedly inhaling it in small doses. Employees in works where water-gas is made, or used, incur the risk both of acute and chronic carbon monoxide poisoning. The recent introduction of water-gas as a domestic illuminant is attended by most serious risks to health, as it is impossible to keep household gas fittings absolutely perfect, and the high percentage of CO present in the gas converts the slightest leakage into an insidious but sure source of chronic poisoning.

<sup>1</sup> *Zeitschr f. physiol. Chemie*, 1888.

<sup>2</sup> *Comptes Rendus de la Société de Biologie*, 1892.

<sup>3</sup> *Arch f. exp. Pathol.*, 1891.

<sup>4</sup> *Loc. cit.*

**Symptoms.**—The earliest symptoms are headache, neuralgic pains, indications of defective nutrition, as anæmia, and loss of flesh, with a sensation of want of breathing power—breathlessness being developed by incommensurate exertion. The more advanced symptoms are those associated with peripheral neuritis, and psychical disturbances. Ross<sup>1</sup> relates the case of a healthy, well-fed, and temperate man employed in making gas, whose business it was to attend to the retorts. After working at this employment for six months, he was pale and anæmic, and was much troubled with shortness of breath, he then began to feel shooting pains in the legs and shoulders, his hands and feet were numb, and he suffered from cramp in the calves of the legs. The fingers were often spastically contracted on the palms, and his gait was “high-stepping.” He improved under treatment. Ruata<sup>2</sup> states that a man suffering from chronic CO poisoning complained of headache, was depressed and became acutely delirious.

### ACETYLENE.

This gas ( $C_2H_2$ ) has an unpleasant odour familiar to all who have been near to a Bunsen burner when the flame has “dropped”, hitherto it has been of little practical interest, but the extension of its use as an illuminating agent increases its importance. Lewin<sup>3</sup> states that an atmosphere containing 1 per cent of acetylene produces deep nœcrosis in dogs with symptoms of asphyxia, more recent experiments, however, tend to show that these results were probably due to impurities, such as  $H_2S$  and  $PH_3$ . Ogier and Brocner<sup>4</sup> came to the conclusion that acetylene is not markedly poisonous. Roseman<sup>5</sup> found that acetylene has a slight narcotic action on animals, but that prolonged exposure to its influence is required to produce actual toxic effects. Frank and Weyl<sup>6</sup> state that acetylene does not possess the poisonous properties ascribed to it, and Gréchant<sup>7</sup> found that it is not poisonous to dogs unless the respired air contains from 40 to 79 per cent of the gas. Mosso and Ottolenghi<sup>8</sup> on the other hand state that acetylene has considerable toxic power, half a litre is mixed with air, so as to constitute 20 per cent of the mixture, proves fatal to dogs. Acetylene does not appear to combine with hæmoglobin, and it is quickly eliminated from the blood, when saturated with the gas, blood yields the spectroscopic reaction of oxyhæmoglobin, and can easily be reduced.

### GASES PRODUCED BY EXPLOSIVES.

In mines in which explosives are used toxic effects have been caused by inhalation of the gases thus generated, the composition of the mixed gases varies with the composition of the explosive, but in all cases the bulk of the gas generated consists of carbon dioxide and nitrogen. Gunpowder in addition yields carbon monoxide in considerable amount, and sulphuretted hydrogen. Nitroglycerine and dynamite also yield a large percentage of carbon monoxide. Gun-cotton yields much the same. Tonite, which is composed of a mixture of equal parts of gun cotton and barium nitrate, produces very little, if any, carbon monoxide. Roburite, which consists of chloro dinitro benzene and ammonium nitrate, when detonated experimentally gives off no CO, the introduction of the last named explosive has been the occasion of a number of investigations as to its alleged injurious effects. Chemical analyses of the gases in the coal mines after the detonation of roburite were separately made by Professor Dixon of Owens College and Professor Bedson of Durham College, the results clearly showed that much less CO (which constitutes the dangerous gas) was given off by roburite than by gunpowder. It will be observed that although experimental detonation of roburite produces no CO, in actual use a small quantity was present, this is partly given off by the fuse when burning and, as suggested by Professor Dixon, is probably also produced by hot CO<sub>2</sub> passing over the coal, in any case the amount is small and the gas is dissipated in a short time. Still, as a very small percentage of CO is deleterious if the air which contains it is long respired, it is important to minimise all risk

<sup>1</sup> *Peripheral Neuritis*, Ross and Bury, 1893

<sup>2</sup> *Gazzetta med di Torino*, 1892

<sup>3</sup> *Lehrb d Toxicologie*, 1885

<sup>4</sup> *Annales d'Hygiène*, 1887

<sup>5</sup> *Arch f exp Path*, 1895

<sup>6</sup> *Nationalzeitung*, 1895

<sup>7</sup> *Comptes Rendus*, 1895

<sup>8</sup> *Riforma Medica*, 1897

of contamination, to this end experts recommend the use of electricity for the purpose of detonating roburite, thus obviating the necessity for a fuse, together with the allowance of sufficient time for dissipation of the products given off by the explosive before the miners resume their work. No evidence was obtained of injurious effects from nitro benzene present in the air after explosion of roburite. The toxic effects of roburite, as such, are dealt with in the chapter on benzene and its derivatives.

## GASES OF WARFARE.

The following account of the gases used by the Germans, and the symptoms they produced, is taken mainly from a pamphlet issued by the Ministry of Pensions in consultation with the Ministry of Health.<sup>1</sup>

**Periods of Gas Warfare.**—(a) The earliest attacks, in April and May, 1915, were carried out by means of drift gas, which was released from cylinders, and was carried down to the trenches by a favouring breeze. The gas at this time employed was pure chlorine. Later and throughout the remainder of this period the drift attacks were carried out with a mixture of chlorine and phosgene. This period lasted until August, 1916, after which drift attacks ceased, as the war became more mobile.

(b) From May, 1915, to July, 1916, in addition to the drift attacks, a certain amount of gas shelling was experienced, but was confined to the use of Lachrymator shells.

(c) From July, 1916, to July, 1917, may be regarded as the period of "Lethal Gas" shelling. In this period shells of all calibres, containing varying mixtures of poisonous gas, the main constituent being phosgene, were fired.

(d) From July, 1917, until the end of the war, gas warfare was characterised by the introduction of various new gases in shells, the main ones being di-chlor-ethyl-sulphide—the so-called "mustard gas"—and the various arsine compounds. These new compounds were used either singly or in mixed bombardments, in which lethal gas, mustard gas, and arsine were employed.

(e) Projector attacks, consisting of drums filled with a large amount of phosgene were also carried out on occasion between December, 1917, and May, 1918.

**Classification of Gases.**—Certain broad distinctions in the action of the various gases may be drawn. These are not absolute, for the action of any gas varies with its concentration. A gas classified as a lung irritant, for example, will, if in sufficient concentration, act as a lachrymator, and a nasal irritant has the power when concentrated of vesicating the skin. Accepting these limitations, however, the following classification has been adopted as affording a working basis, as regards the gases known to have been employed by the enemy—

### *Lung Irritants—*

Chlorine,  
Phosgene,  
Chlor-methyl-chloroformate,  
Tri-chlor-methyl-chloroformate,  
Chloropierin,  
Phenyl-carbylamine-chloride,

### *Formula*

$\text{Cl}_2$   
 $\text{COCl}_2$   
 $\text{CH}_2\text{Cl COOCl}$   
 $\text{CCl}_3 \text{ COOCl}$   
 $\text{CCl}_3 \text{ NO}_2$   
 $\text{C}_6\text{H}_5\text{N C Cl}_2$

### *Nasal Irritants—*

Di-phenyl-chlor-arsine,  
Ethyl-di-chlor-arsine

$(\text{C}_6\text{H}_5)_2 \text{As Cl}$

<sup>1</sup> "Notes and Suggestion on (1) Dysentery, (2) Trench Fever, and (3) Gas Poisoning and its Sequels," 1920.

*Lachrymators*—

Benzyl Bromide,

Xylol Bromide,

Brom-acetone,

Mono-brom-methyl-ethyl-ketone,

Di-brom-methyl-ethyl-ketone,

 $\text{C}_6\text{H}_5\text{CH}_2\text{Br}$  $\text{C}_6\text{H}_4\text{CH}_2\text{CH}_2\text{Br}$  $\text{CH}_3\text{Br CO CH}_3$  $\text{CH}_2\text{Br CO CH}_2\text{CH}_3$  $\text{CH}_3\text{CO CHBr CH}_2\text{Br}$ *Vesicant*—

Di-chlor-ethyl-sulphide (Mustard Gas),

 $(\text{CH}_2\text{Cl CH}_2)_2\text{S}$ 

**Lung Irritants**, of which phosgene may be taken as a type, act as direct irritants to the pulmonary alveoli—the upper respiratory tract escaping except in high concentration. The result is pulmonary oedema with rupture of alveolar walls, patchy emphysema, and thrombosis of capillaries—the latter most marked at the vulnerable infundibulo-alveolar junction. Further, there rapidly develops a high concentration of blood, to the production of which several factors contribute, among them being the loss of fluid from pulmonary oedema, and the shock element. These are the primary effects, and they are followed, in pathological sequence as might be expected by secondary infections with the common bacteria of the respiratory passages in cases where the initial inflammatory changes have been severe.

Within three days recovery, in the non-fatal cases, commences, and unless secondary infections develop, is rapid, so that convalescence is approaching by the end of a week. One of the early valuable signs of approaching convalescence is bradycardia, which may be seen even after extremely mild gassing. Where convalescence is more prolonged debility, gastric disturbance, and pain in the chest are the most frequent complaints.

Should secondary complications develop, they present the ordinary features of pleurisy, bronchitis, or broncho-pneumonia.

The proportion of men who suffered from permanent or prolonged after-effects is small. Broncho-pneumonia and pleurisy may have left permanent lesions in the form of patches of fibrosis, pleural adhesions, etc. The extreme lack of oxygen during the acute stage of pulmonary oedema has led in some cases to a cardiac condition with precordial pain, dyspnoea, exhaustion, and persistent tachycardia, after exercise, comprising the symptoms usually referred to as D.A.H. or the “effort syndrome.” Another type is characterised by recurring attacks of nocturnal dyspnoea, in which the patient is aroused from sleep by the sudden onset of difficult breathing, the respiration being shallow and rapid. In rare cases, chronic albuminuria has been observed.

**Lachrymators.**—With concentration obtainable in the field, the main action of the vapours of these substances is to cause profuse lachrymation and smarting of the eyes, which may be so extreme as to prevent the eyes being opened. There may be a transient condition of conjunctivitis and oedema of the eyelids. The condition rapidly clears up, and as there are no toxic after-effects, further consideration is not required.

**Nasal Irritants.**—The action of the arsines is essentially that of an irritant to the upper respiratory passages, and the oesophagus and stomach. The liquid ethyldichlorarsine applied to the skin will cause blisters, but this is not a condition likely to be met with frequently in the field. The arsines also have an influence on the central nervous system, causing either mental irritability or more frequently drowsiness. Further evidence of this involvement is at times

seen in sensory changes varying from numbness or tingling of the finger tips to a complete anæsthesia, usually of the glove-and-stocking type, and involving all forms of sensibility. How far the nervous symptoms are due to a neurosis, or how far they are based on a definite organic lesion, presumably a transient toxæmia of cell elements in the central nervous system, has not been determined.

**Diphenylchlorarsine.**—Almost immediately the following symptoms occur.—Pain in the nose and sneezing, and frequently there is watering or irritation of the eyes. These are rapidly followed by tracheal and pharyngeal pain with a feeling of constriction of the throat. Then nausea and vomiting occur, and there is a feeling of swelling of the face and throat, but no evidence of actual swelling or œdema. In course of time these symptoms abate, and vomiting is replaced by gastric discomfort. The last symptom to disappear is the pain in the nose and forehead (24-48 hours). After 24 hours the patients are practically recovered, and they only complain of weakness. Usually, at first, mental irritability is present, or there may be drowsiness.

**Ethylidichlorarsine.**—The symptoms are very similar to the foregoing. There may, in addition, be a mild bronchial irritation lasting for a few days. The nervous symptoms are more pronounced and more frequent. No fatal case following arsine poisoning has been reported. Beyond relieving pain by chloroform or opium, no treatment was found necessary.

**Vesicants.**—These are essentially chemical irritants, causing burns which subsequently become infected. Lasting results of vesicants include scarring of the skin from burns, permanent lesions of the eyes, such as corneal ulceration, keratitis, and nebulae, ulceration or thickening of the vocal chords leading to impairment of movement and chronic bronchitis, localised fibrosis of the lungs, and pleural adhesions.

Various functional symptoms, such as photobia, aphonia, and vomiting, have persisted for a long time after gas poisoning.

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## CHAPTER XXXIII

### POISONING BY COMPOUNDS OF CARBON, HYDROCYANIC ACID, AND CYANIDES.

**Hydrocyanic Acid (HCN)**, or prussic acid, in its commercial form contains from 2 to 5 per cent of the anhydrous acid. It has a penetrating odour, but when smelt in very dilute solution it produces a bitter taste at the back of the tongue, rather than olfactory sensation. It is feebly acid, slightly reddening litmus-paper. Unless hermetically sealed up and kept in the dark, it loses strength more rapidly than is generally supposed.

**Oil of bitter almonds**, used as a flavouring adjunct, contains a varying percentage of anhydrous hydrocyanic acid, in the crude oil amounting to from 5 to 15 per cent. **Cherry-laurel water** contains about 0.1 per cent.

**Potassium Cyanide (KCN)**, largely used in photography and in electroplating, is a salt which has a strong alkaline reaction. Hydrocyanic acid is displaced from KCN by carbon dioxide; therefore the percentage of acid diminishes by keeping, the "cyanide" of commerce usually contains some potassium carbonate, and from absorption of oxygen tends to be converted into

cyanate Potassium cyanide yields the odour of hydrocyanic acid without the addition of a displacing acid Several other cyanides are poisons, but are rarely used as such

Many plants of the natural order *Rosaceæ*, especially of the sub-orders *Prunææ* and *Pomeææ*, contain a crystalline substance—**Amygdalin**—which is capable of yielding hydrocyanic acid Amygdalin itself is not poisonous, but when it is subjected to the action of a natural ferment—emulsin—which is found along with it in the fruit or leaves of the above-named sub-orders, it is decomposed into oils of bitter almonds, glucose, and hydrocyanic acid, the same result is produced when amygdalin is boiled with dilute acid In cases of criminal poisoning with hydrocyanic acid endeavours have been made to utilise the occurrence of hydrocyanic acid-forming substances in edible fruit to account for its presence in the human body in amount sufficient to cause death bitter almonds excepted, it is very improbable that a sufficient quantity of such fruit could be eaten as to introduce a fatal dose of hydrocyanic acid

**Symptoms.**—When a fatal dose of hydrocyanic acid is swallowed the symptoms usually appear within a few seconds, they may be delayed for thirty or forty seconds, and exceptionally for a little over a minute, when thus delayed, the victim may be able in the interval to walk and speak When animals are poisoned with hydrocyanic acid they almost invariably give utterance to a spasmodic cry, this symptom is frequently absent in human beings After a few gasps the patient becomes insensible, and, if in the upright posture when the poison is swallowed, he falls to the ground The surface is cold, the face usually pallid, the eyes are open and staring, and the pupils are dilated and insensible to light The breathing is laboured, irregular, and gasping, the breaths being drawn at longer and longer intervals, after apparently final cessation, conclusive efforts at respiration may occur with prolonged intervals between Tetanic spasms affecting the muscles of the jaws and limbs usually occur in the early stage, followed by complete relaxation of all the muscles The pulse is almost or quite imperceptible at the wrist, if it can be felt it is found to be exceedingly rapid—120 or more in the minute—small and irregular, if death does not take place immediately the pulse may become slower towards the end The lips are frequently covered with froth, vomiting, involuntary micturition and movement of the bowels are not infrequent Death usually takes place within five or ten minutes, it may occur almost immediately after the poison is swallowed, or, when a minimum lethal dose is taken, the fatal issue may be delayed for an hour and even longer Stevenson<sup>1</sup> records a case in which death did not take place for an hour and a quarter If life is prolonged over half an hour there is considerable chance of recovery

The mode in which hydrocyanic acid causes death has been the subject of much discussion The earlier opinion, founded on the exceeding rapidity of its action, was that it paralysed the central nervous system (Preyer)<sup>2</sup> This explanation has recently been upheld by Corn and Ansaux,<sup>3</sup> who believe that the hydrocyanic acid paralyses the vaso motor centre, and also by Masius,<sup>4</sup> and others, who attribute death to paralysis of the medullary centres by which respiration and tissue-oxidation are suddenly arrested Another view, originally propounded by Schonbein,<sup>5</sup> is that death is produced by arrest of internal respiration, without the intervention of the nervous system It has been found experimentally by Gaethgens<sup>6</sup> that in poisoning by hydrocyanic acid less carbon dioxide is excreted and

<sup>1</sup> *Guy's Hosp Reps*, 1869

<sup>2</sup> *Die Blausaure*, 1868-70

<sup>3</sup> *Bull de l'Acad Belgique*, 1894

<sup>4</sup> *La Semaine Med*, 1894

<sup>5</sup> *Zeitschr f Biologie*, Bd. III

<sup>6</sup> *Hoppe-Seyler Med Chem Untersuch.*

less oxygen taken up than in the normal, the imperfect oxidation which ensues is shown by the presence, in the blood, of lactic acid as a substitution product for carbon dioxide (Zillessen).<sup>1</sup> Geppert<sup>2</sup> explains the arrest of internal respiration on the hypothesis that hydrocyanic acid deprives the tissues of their power to take up oxygen, according to this view the oxygen of the blood being neglected by the tissues accumulates until the whole of the blood—venous as well as arterial—assumes a bright red colour, asphyxia thus takes place in the presence of excess of oxygen. Schonbein attributes the arrest of internal respiration to the action of hydrocyanic acid on the blood. Kobert,<sup>3</sup> whilst accepting Geppert's theory, believes that HCN also kills the protoplasm of the red corpuscles, rendering them functionless as oxygen carriers, and further, that it directly attacks the nervous system. He holds that HCN forms a definite compound with methæmoglobin—cyanmethæmoglobin—which is bright red in colour and yields a spectrum resembling that of reduced hæmoglobin, it possesses considerable stability and resists the reducing influence of the tissues so as to be capable of recognition eight days after death in the blood of an individual poisoned with hydrocyanic acid, it is not affected by ammonium sulphide, nor does the passage of a current of air through it dissociate the HCN. Kobert assumes that after death HCN combines with methæmoglobin, and that the resultant cyanmethæmoglobin by reason of its bright red colour determines the peculiar tint of the post mortem stains and also that of the mucous membrane of the stomach. Szigeti<sup>4</sup> regards cyanmethæmoglobin as being identical with cyanhæmatin, a substance discovered by Hoppe-Serler.<sup>5</sup>

**Fatal Dose.**—The smallest recorded fatal dose was half a drachm of the B P acid—equal to 0.6 grain of anhydrous acid—which caused death in one hour and twenty minutes, Garstang,<sup>6</sup> who records the case, states that hydrocyanic acid was inadvertently substituted for hydrochloric acid in dispensing a prescription, therefore the amount taken could be accurately ascertained. The largest dose followed by recovery was half an ounce of medicinal acid, equal to 4.8 grains of anhydrous acid, the case is described by Shively.<sup>7</sup>—A student of pharmacy was found insensible, the pupils being widely dilated, the right more so than the left, respiration was laboured and there was extreme dyspnoea, with coldness of the surface, but without lividity or cyanosis, the face having a rosy tint. The pulse was 86 per minute, but rapidly went up to 112, and was thready and irregular, the temperature in the rectum was 97.5° F. Trismus and rigidity of the limbs were present for a short time. The stomach-tube was used, and alcohol, camphorated ether, and atropine sulphate were administered, the two latter subcutaneously, the alcohol by mouth and rectum, faradisation of the phrenics was also resorted to. The urine contained albumin, crystals of calcium oxalate, and gave a precipitate with ferric and ferrous salts. On recovery the patient explained that the first sensation he experienced after swallowing the poison was numbness of the lips, which was quickly followed by shortness of breath and loss of consciousness. In another case<sup>8</sup> recovery took place after two drachms—about 2.5 grains of anhydrous acid. A woman swallowed the dose by mistake, after which she rushed upstairs to her master, who was a medical man, told him what she had done, and then fell down insensible, the stomach-tube was used immediately, and apomorphine was given, followed by electricity and artificial respiration, the treatment being attended with success. This case is remarkable, not only for recovery after a dose more than double that which is usually fatal, but also for the actions performed by the patient in the interval that elapsed between swallowing the poison and the occurrence of insensibility, the power of moving

<sup>1</sup> *Zeitschr f physiol Chemie*, 1891

<sup>2</sup> *Zeitschr f. klin Med*, 1889

<sup>3</sup> *Ueber Cyanmethæmoglobin und den Nachweis der Blausäure*, 1891.

<sup>4</sup> *Vierteljahrsschr f ger Med*, 1893

<sup>5</sup> *Med chem Untersuch*

<sup>6</sup> *The Lancet*, 1888

<sup>7</sup> *Internat Journ Med Sciences*, 1890

<sup>8</sup> *Brit Med Journ*, 1890

and speaking after swallowing fatal doses has been observed in many cases, but not often, if ever, to such a degree

In cases of fatal poisoning by very small doses, and of recovery after very large ones, a doubt exists as to the percentage of anhydrous acid present in the solution swallowed. Examination of specimens of dilute acid obtained from several sources showed that some were above and some below the standard strength. It may be accepted that an amount of dilute acid which contains one grain of anhydrous acid constitutes a fatal dose. The action of a fatal dose of hydrocyanic acid is not diminished by dilution though it may be slightly delayed.

**Treatment.**—Immediate evacuation of the stomach with stomach-tube, or emetics. Jena recommends that three drachms of a 1 in 1,000 solution of adrenalin diluted should first be given in order to delay the absorption of the poison and give more time for subsequent treatment. After repeated washing out a further small dose of adrenalin may be left in the stomach. Zinc sulphate or mustard may be given by the mouth, or apomorphine subcutaneously, then artificial respiration, faradisation of the phrenics and the diaphragm, external warmth and friction, subcutaneous injections of ether, brandy by the mouth or rectum, and if the surface is not cold, effusion with cold water. If effusion is resorted to it should be intermittent either vigorous friction or hot applications being used in the intervals. Subcutaneous injections of atropine have been recommended on theoretical grounds—stimulation of the respiratory centre the advantage is doubtful. Chemical antidotes, practically speaking, are useless first, because all the poison should be removed from the stomach, and not left for chemical neutralisation, and second, because the action of the poison is far too rapid to permit of the effective administration of an antidote that requires special ingredients and time to prepare it. Recently two antidotes have been proposed which, it is stated, will not only neutralise any HCN that may be present in the stomach, but, when injected subcutaneously, will also act upon that which has been absorbed. Antal<sup>1</sup> recommends a 0.5 to 1 per cent solution of cobalt nitrate, and Lang<sup>2</sup> a 5 to 10 per cent solution of sodium thiosulphate. Twenty minims or more of one or other of these solutions may be repeatedly injected under the skin, the resulting combinations with HCN—cobalt cyanide and sodium sulphocyanide respectively—are harmless. Martin and O'Brien propose as an antidote 1 oz of ferrous sulphate solution of 23 per cent strength, 1 oz of KOH solution 5 per cent strength, and 2 grains of magnesia. These to be mixed and taken immediately, the object being to form the comparatively innocuous Prussian blue.

**Post-mortem Appearances**—**External.**—The eyes, with dilated pupils, may be prominent and bright, the fingers and the jaws clenched, the lips may be covered with froth, and the post-mortem stains are frequently pink or light red. **Internal.**—On opening the abdomen the odour of HCN may be perceived. If not, it may sometimes be recognised on removing the calvarium. The blood may be bright red in colour, but it is often dark, it is almost always fluid. The mucous membrane of the stomach may be brilliant red, due to the colour of the blood, all the mucous surfaces and even the muscles may show more or less a like tendency. The right side of the heart is usually distended with blood.

**Potassium Cyanide** produces symptoms like those of HCN, with the addition of more or less local action on the mucous membrane of the mouth and stomach.

<sup>1</sup> *Experimentelle Untersuchungen zur Therapie der Cyanvergiftungen*, 1895.

<sup>2</sup> *Arch. exp. Path.*, 1895.



The lips and mouth may be eroded or the mucous membrane softened and capable of being easily detached. The mucous membrane of the stomach has been found partially or wholly of a bright red, deeply injected, thickened, softened, and even eroded, its surface may be covered with blood-stained mucus. The stomach-contents will be alkaline in reaction until all the poison is evacuated.

**Fatal Dose.**—Death has resulted from five grains, and recovery after about forty grains were taken in the solid form.

**Oil of Bitter Almonds.**—In one case<sup>1</sup> a teaspoonful of the crude oil, which was afterwards found to contain 3·4 per cent of HCN, caused the death of a girl in an hour and three-quarters, the dose taken contained about two grains of anhydrous HCN. The contents of the stomach yielded HCN fourteen days after death. It has been frequently noticed that the odour of HCN in the dead body lasts longer when received into the system in the form of oil of bitter almonds than as a solution of pure hydrocyanic acid.

Baker<sup>2</sup> records the case of a man who ate two handfuls of **bitter almonds**, he went to his work, but soon after became insensible, with all the symptoms of poisoning by hydrocyanic acid, prompt use of the stomach-pump and active treatment led to recovery. The contents of the stomach yielded the reactions of HCN.

**Chemical Analysis- Tests.**—A drop of a solution of silver nitrate placed on the concave surface of a watch-glass, and inverted immediately over a substance which contains free hydrocyanic acid, becomes milky from the formation of silver cyanide. If the amount of HCN is small, the milky appearance first manifests itself as a white line around the margin, which gradually spreads over the whole of the drop, when slowly formed the deposit, examined under the microscope, is seen to consist of delicate acicular or prismatic crystals, if rapidly formed a crystalline mass is produced without distinctive appearance. Silver cyanide is soluble in hot concentrated nitric acid. A drop of a solution of potassium hydroxide substituted for the silver solution, and exposed as before to the vapour of HCN, undergoes no visible change on being allowed to remain for a few minutes. A drop of a solution of ferrous sulphate subsequently added to the potash produces a greenish-grey precipitate, which becomes blue (Prussian blue) on the addition of dilute hydrochloric acid. A drop of ammonium sulphide on a watch-glass inverted as before, allowed to remain two or three minutes, and then evaporated to dryness at a gentle heat, yields a blood red colour (ferric sulphocyanide) on being touched with a glass rod dipped in a solution of ferric chloride, the colour disappears on the addition of a drop or two of a solution of mercuric chloride.

If the temperature is low, it may be necessary to warm slightly the substance tested. In order to liberate HCN from potassium cyanide, the addition of tartaric acid is necessary, in sufficient amount to produce an acid reaction.

Robert directs attention to the property possessed by HCN of preventing and of abolishing the reaction of iodine with starch, and suggests that it constitutes an exceedingly delicate test, which may be performed in two ways.—If a little very dilute cold solution of starch (prepared by boiling) with potassium iodide, is divided between two test tubes, to one of which a minute quantity of HCN is added, on dropping into each tube some aqueous solution of peroxide of hydrogen the contents of that to which HCN was added remain unchanged, those of the other become blue. The second way is to tinge a little dilute solution of starch with iodine, on the addition of a minute amount of HCN the colour is discharged. A few drops of a distillate obtained from the blood of an individual poisoned

<sup>1</sup> Reg. v. Timins (Maidstone Assizes, 1883)

<sup>2</sup> *Brit. Med. Journ.*, 1881.

with HCN may be thus tested. The same reaction, however, is produced by  $H_2S$  and some other substances. Although HCN is very liable to undergo decomposition, it has been detected in the tissues four months after death.

**Quantitative Estimation**—The organic admixture if necessary should be rendered acid with tartaric acid, and distilled over a water bath. It is difficult, if not impossible, to get all the HCN over, as some of it will probably be decomposed. The amount of cyanogen present in the distillate is best estimated by titration with a standard solution of silver nitrate. If preferred the distillate may be acidulated with nitric acid, and precipitated with silver nitrate, the precipitate is washed, dried, and weighed—100 parts correspond to 20.15 of anhydrous hydrocyanic acid.

## CARBON COMPOUNDS—FATTY GROUP.

### ALCOHOL.

**Acute** poisoning with alcohol ( $C_2H_6O$ ) is the only form that it will be necessary to describe, the chief medico-legal interest being centred in the diagnosis. As a rule the lesser degrees of alcoholic intoxication are easily recognised, the difficulty arises when a profound comatose stage is reached, and no history is forthcoming to aid diagnosis—as, for example, when a medical practitioner is called upon by the police to determine the condition of a man found in a state of insensibility in the street or other public place, in order to arrive at a reliable conclusion in such instances, method is necessary.

The first consideration is—What toxic and pathological conditions may be mistaken for profound alcoholic intoxication? Among these are **Cerebral lesion**, as embolism, hæmorrhage into the pons, internal capsule, or cortex, **poisoning by opium**, chloral hydrate or other narcotics, **mechanical injury**, as a blow or fall on the head, **diabetic**, or **uræmic coma**, **post-epileptic coma**, and some forms of **hysterical** or non-organic **neuroses**.

An initial difficulty presents itself, two of these conditions may be combined. A drunken man may have received a violent blow on the head, or he may be the victim of cerebral hæmorrhage due to rupture of a vessel, apart from external violence. First, as to the pupils, if dilated they suggest alcohol, if contracted, opium, or hæmorrhage into the pons. In hæmorrhage into the pons the temperature is frequently elevated ( $103.5^\circ$ ) from the first, in the stage of opium poisoning that corresponds the temperature is subnormal. If one pupil is dilated and the other contracted, or normal, some intracranial lesion is probable. Conjugate deviation of the eyes suggests hemiplegia. Note if there is flapping of one cheek, and if the arm and leg are equally limp on both sides, lift up alternately the arm and the leg on one side and let them fall, and then repeat the process on the other side, comparing the results, in hemiplegia, unless there is initial rigidity, the limbs on the paralysed side will fall more like inanimate objects than those on the unaffected side. Pinching the skin will sometimes evoke movements that reveal the presence or absence of hemiplegia, the plantar reflex may be tried with the same object. The great toe is extended on the hemiplegic side and is flexed on the sound side (Babinski). Bilateral inequality of any of the reflexes is suggestive of organic trouble. If by shouting into his ear the patient can be sufficiently roused to give his name, occupation, and address, the stupor is not likely to be due to organic lesion, it may result from opium in the early stage, or from alcohol. The odour of alcohol in the breath is of little significance, since brandy is often given as a remedy to patients found insensible, its absence, however, eliminates alcoholic poisoning from the possible causes of insensibility. If opium as such, or in

the form of tincture, is the cause of insensibility, its odour may possibly be perceived in the patient's breath. Examine the head for indications of bruising, or cuts of the scalp, or of fracture of the skull, look for signs of hæmorrhage from the nostrils or ears. It is to be remembered that comparatively slight blows, or falls on the head, may produce effusion of blood into the arachnoid, which often does not cause any obvious symptoms for some time after the injury has been inflicted, this is especially the case when the condition is masked by alcoholic intoxication.

In **alcoholic coma** the face may either be flushed or pale, the pupils either contracted or dilated—frequently contracted at first and afterwards dilated. The stertor is not usually so marked as in apoplexy, unless a fatal issue is imminent. If on using the stomach-tube a large quantity of strongly alcoholic fluid is removed, the diagnosis of alcoholic poisoning is strengthened. The severest form of acute alcoholic poisoning occurs when the patient has obtained access to an unlimited amount of spirit (as by boring a hole in a whisky-cask), and has swallowed enormous doses of it undiluted, the symptoms quickly develop into those of coma of a profound type, which, in the absence of any evidence of a focal lesion, cannot be distinguished from that due to apoplexy. The presence of alcohol in the urine of those who have partaken freely of it may be ascertained by the potassium dichromate test subsequently described.

In **uræmia** the pupils will be contracted, and there will be recurrent convulsions, the temperature will be subnormal. The presence of albumin in the urine is not of much diagnostic value, as it is frequently found in apoplexy. **Post-epileptic** coma is chiefly met with in young people, it resembles profound sleep rather than true coma, the tongue should be examined for injuries inflicted by the teeth. **Diabetic coma** has been mistaken for drunkenness, not only in the comatose stage, but also in the preceding stage of excitement, which strongly resembles that due to alcohol. The diagnostic signs are a peculiar odour of the breath like that of American apples with sugar and probably acetone and diacetic acid in the urine, the respirations are slow and sighing, and the temperature is considerably below the normal. Exceptionally there may be no odour of acetone. Unconsciousness, due to **hysteria**, may resemble that due to meningitis. The diagnostic signs are the age and sex, the absence of indications of cerebral lesion, with probably normal temperature, pulse, and condition of the skin.

**Treatment.**—The stomach-tube is to be resorted to, or in its absence an emetic. Faradisation, or flicking the patient with a wet towel, compelling him to attempt to walk with the aid of a man on either side of him, cold douche, with alternate friction, and the administration of hot coffee are the means to be used for promoting return to consciousness. If there is doubt as to the cause of coma, treat the case as possibly due to cerebral lesion, and keep the patient under observation until the diagnosis is established.

Alcohol is eliminated by the kidneys and the lungs.

**Post-mortem Appearances.**—The most characteristic appearances of acute alcoholic poisoning are found in those who have died shortly after swallowing large doses of alcohol. Cadaveric rigidity is usually well marked, and sometimes lasts for several days, putrefactive changes advance slowly. The odour of the spirit partaken of is perceived on opening the stomach, unless it was well washed out before death, the abdominal, thoracic, and cranial cavities yield similar evidence. The mucous membrane of the stomach is sometimes injected and bright red in colour, at others it is pale, with or without isolated reddened spots. The right heart and the veins are usually filled with dark fluid blood,

and the lungs are hyperæmic, either throughout or at the posterior part of the lower lobes. The bladder generally contains a large quantity of urine. The cerebral vessels as a rule are well filled with blood, and there may be extravasations in the membranes or in the substance of the brain.

If the deceased was an habitual drunkard, the ordinary pathological changes due to chronic alcoholism will be present in addition to the appearances above described.

**Chemical Analysis**—Alcohol may be separated from organic admixture by distillation at a gentle heat. If the substance to be distilled is strongly acid, sodium carbonate should first be added until the reaction is neutral. redistillation may be necessary, along with rectification by lime or potassium carbonate.

**Tests.**—On heating some of the distillate in a test-tube with a few drops of a solution of potassium dichromate and a little sulphuric acid, the colour changes from yellow to green, and the odour of aldehyde is given off. The odour of acetic ether may be produced by heating some of the distillate with an equal volume of sulphuric acid and an acetate. To a little of the distillate, in a test-tube, add ten or twelve drops of a strong aqueous solution of iodine (dissolved with the aid of potassium iodide), mix and add a solution of potassium hydroxide until the colour is reduced to a light yellow. On gently heating, the clear fluid becomes cloudy, owing to the formation of iodoform, which may be recognised by its odour, and, if slowly formed, by the appearance under the microscope of the crystals which are presently deposited, they take the form either of rosettes, or small hexagonal plates like crystals of cystin. If the solution of alcohol is weak, boil it with the iodine solution for several seconds and then rapidly cool the test-tube in a stream of cold water, when the clear liquid becomes turbid from the precipitation of iodoform. It is to be remembered that other substances besides alcohol—as aldehyde and acetone—give the iodoform reaction.

**Quantitative estimation**, as a rule, is not feasible, or at least would not convey any idea of the amount of alcohol swallowed.

**Methyl Alcohol** when pure closely resembles ethyl alcohol in physical properties. In the crude state, as wood naphtha, it is nauseous to both nostrils and palate, and is used to render ethyl alcohol, for trade purposes, unpalatable. This compound is known as methylated spirit, and notwithstanding its repulsive odour it has been drunk to such an extent as to cause the Excise authorities to substitute other methods of preventing the abuse of alcohol only intended for manufacturing purposes. The symptoms produced by methylated spirit resemble those due to ethyl alcohol, of which the former largely consists. A prominent additional symptom caused by methyl alcohol is a form of amblyopia, the condition may consist in simple temporary derangement of the ocular circulation, or it may go on to acute retrobulbar optic neuritis. Cases of this kind have been reported by Nagel,<sup>1</sup> Stromberg,<sup>2</sup> and others.

**Paraldehyde**  $[(C_2H_5O)_2]$ , when taken in excess, has, in a few instances, produced toxic symptoms, and one instance is recorded in which death took place after two ounces. Mackenzie<sup>3</sup> relates a case in which three and a half ounces were swallowed, and produced, for thirty-four hours, a condition resembling chloroform narcosis, strychnine was administered hypodermically, and the patient recovered. Fornace and Quarrelli<sup>4</sup> record the case of a man who, at the age of forty-two, began to take paraldehyde in doses of two grammes daily for insomnia. During the next five years this was gradually raised to fifteen grammes a day. Alternate fits of excitement and depression developed, with tremor of the hands and disturbance of speech. This led to larger doses, and in one week he took 500 grammes. Eventually he took 100 grammes of the drug and then went to hospital, where he was admitted partially unconscious. For eight days he was delirious, the temperature at the

<sup>1</sup> *Journ. Amer. Med. Assoc.*, 1905.

<sup>2</sup> *St. Petersburg med. Wochenschr.*, 1904.

<sup>3</sup> *Brit. Med. Journ.*, 1891.

<sup>4</sup> *Berl. klin. Woch.*, 1912.

height of the delirium being  $104^{\circ}$  and the pulse 134. Perspiration was profuse. The delirium gradually subsided, and on the twentieth day he returned to work. Opium in the form of pantopon was found to be the most beneficial treatment.

**Formic Aldehyde** ( $\text{CH}_3\text{O}$ ) has recently come into general use as a disinfectant and preservative of organic substances. For these purposes the commercial preparation called **Formalin**, which is usually a 40 per cent solution of formic aldehyde, is used. Although regarded as being but feebly poisonous, it has produced severe toxic symptoms, and even death. Zorn<sup>1</sup> records the case of a man aged forty-four, who drank about half a fluid ounce of formalin (40 per cent). Retching and vomiting occurred, the pulse was 126, small and regular, and the respirations were 44 to the minute. The lips and extremities were cyanotic. No urine was excreted for twenty-four hours, and the first subsequently voided contained albumin, but no blood nor sugar. The patient was dizzy and the gait was uncertain. The motions were passed with much straining, they contained mucus but no blood. Recovery took place in a few days. In a case related by Kluber,<sup>2</sup> a man drank a mouthful of commercial formalin. He became unconscious, with cold clammy surface, as is the case after an excessively large dose of alcohol. The respiration rate was accelerated, but the pulse and temperature remained normal. There was reddening of the conjunctiva and of the mucous membrane of the throat. There was no vomiting. The urine was completely suppressed for nineteen hours, that subsequently passed contained formic acid until the second day, but no albumin or sugar. Complete recovery took place. Andre<sup>3</sup> saw a case in which a tablespoonful of formalin (40 per cent) produced immediate violent pain in the stomach, great anxiety, and distension of the intestines with gas. The symptoms were relieved by the prompt administration of liquor ammoniæ acetatis, which decomposes the formalin with the liberation of free acetic acid, which in its turn is neutralised by the further administration of alkalies, or magnesia. Bock<sup>4</sup> relates a fatal case in which a man aged twenty-six drank from one to three ounces of a 4 per cent solution of formic aldehyde. Immediate pain in the stomach and vomiting of blood-stained matter, which had the pungent odour of formalin, occurred. The patient gradually became weaker, and died in thirty-two hours from heart failure. Post-mortem, the upper two-thirds of the œsophagus was slightly inflamed, the cardiac end of the stomach was deeply inflamed, and the stomach wall was necrotic, dark and tough, and it cut like old leather. The valvulæ conniventes of the duodenum were inflamed. Watt<sup>5</sup> records the case of a man of sixty-three who died in less than four hours after taking an ounce of formalin containing 34 per cent of formaldehyde. Inhalation of the vapour of formalin has produced toxic effects.

**Tests.**—Formalin reduces ammonio nitrate of silver. If a weak aqueous solution of aniline is added to a solution containing formalin a white precipitate of anhydroformaldehyde aniline is produced. Put a little salicylic acid into a dry test tube and dissolve it in two or three cubic centimetres of strong sulphuric acid. On the addition of a drop of formalin the solution becomes deep red in colour. Dissolve half an inch deep of resorcinol in a test tube, in two or three cubic centimetres of solution of potash. Add a little formalin and boil, the original yellow colour gradually changes to red. If a few drops of a saturated alcoholic solution of gallic acid are added to a solution which contains formic aldehyde, and the mixture is trickled down a test tube on to some concentrated sulphuric acid, a green or bluish ring forms at the junction of the two layers.

**Ether** ( $\text{C}_2\text{H}_5\text{O}$ ) is of little interest as a poison, when swallowed in the liquid form it causes symptoms resembling those due to alcohol. In parts of Ireland it is habitually used for intoxicating purposes, Hart<sup>6</sup> gives an exhaustive account of the subject, showing that the habit is wide spread. Two to four drachms is the usual intoxicating dose, but those used to drinking it can take an ounce or more. Cohn<sup>7</sup> states that in Lithuania, where ether-drinking is common, those accustomed to the habit will drink a quarter of a litre—nearly nine fluid ounces—at once.

**Amyl Alcohol** ( $\text{C}_5\text{H}_{11}\text{O}$ ), or fusel oil, is formed in the manufacture of ethyl alcohol from grain, potatoes, must of grapes, and other sources. Crude fusel oil is an admixture of ethyl, propyl, and butyl alcohols and their ethers, the principal component being amyl alcohol, fusel oil prepared from potatoes consists of about equal parts of ethyl and amyl alcohols, with traces of some of the other alcohols.

Amyl alcohol is of lighter specific gravity than water, with which it is but slightly miscible. It is an oily, colourless liquid, having an acid taste and a peculiar odour, its vapour is very irritating to the respiratory organs, producing a sense of suffocation, accompanied by coughing, and if inhaled for a short time it causes headache.

<sup>1</sup> *Munchener med Wochenschr*, 1900

<sup>6</sup> *Fort Wayne Med Journ -Mag*, 1899

<sup>2</sup> *Munchener med Wochenschr*, 1900

<sup>7</sup> *Brit. Med Journ*, 1912

<sup>3</sup> *Journ de Pharm*, 1899

<sup>4</sup> *Brit Med Journ*, 1890

<sup>5</sup> *Wochenschr f ger Med*, 1898

In a case recorded by Ord,<sup>1</sup> the **symptoms** of acute poisoning by fusel oil were as follows :—A man, aged sixty four, drank about half a pint of fusel oil, which was afterwards found to consist of equal parts of amyl and ethyl alcohols, he felt no ill effects for four and a half hours, when he became unconscious. The muscles were slightly rigid, the teeth were tightly clenched, the face was flushed, but the surface was cold, the respirations were shallow and slow, the pulse was only just detectable at the wrist, the pupils were small, and acted feebly to light. The breath had an odour resembling amyl nitrite, or essence of pears, later on the breathing ceased, necessitating artificial respiration several times, the pulse continuing in the meanwhile. The urine contained both amyl and ethyl alcohols. Recovery took place.

Swain<sup>2</sup> records a fatal case of poisoning by "faints," a refuse after distillation of spirit from potatoes, consisting of a mixture of amyl, propyl, and other alcohols. The mucous membrane of the stomach was soft and thick, and the organ contained a grumous fluid tinged with blood. The odour of amyl nitrite, but sweeter, was perceived on opening the body, and the fluid found in the ventricles of the brain was also odorous. No cirrhotic changes were found in the liver or kidneys, although the patient had frequently indulged in raw amyl alcohol.

**Test**—By distillation with potassium acetate and sulphuric acid, amyl acetate is produced, which is known commercially as essence of jargonelle pears, it may be recognised by its odour.

**Amyl Nitrite** ( $C_5H_{11}NO_2$ )—A case, interesting in more than one respect, is recorded by Rosen.<sup>3</sup> A student, aged twenty two, who was subject to epileptic seizures, had some amyl nitrite given to him for treatment by inhalation. On one occasion, feeling an attack imminent, he got the bottle of amyl nitrite in order to inhale some, he had a fit, and on recovering consciousness, experienced sensations which convinced him that he had drunk some of the fluid during a state of epileptic automatism. Eructations and vomiting occurred. When seen the face was pale, the lips were bloodless, the respirations quiet, and the pulse 110 per minute, he had pain in the head, was much depressed, felt a burning sensation in the throat, and oppression in the region of the stomach, the mucous membrane touched by the poison was slightly eroded, gastric catarrh ensued, with ultimate recovery. The amount swallowed was from twelve to fifteen grammes.

## NITROGLYCERINE.

**Nitroglycerine** [ $C_3H_5(NO_2)_3O_8$ ] is an oily liquid which detonates violently on percussion; it is very slightly soluble in water, and soluble in alcohol and ether. It produces the physiological effects of a nitrite in a powerful degree.—The arteries relax, causing a sensation of fulness and throbbing in the head, frequently accompanied by violent pain, the action of the heart is quickened, and the blood tension lowered, paralysis, both motor and sensory, occurs, and death from respiratory paralysis. Nitroglycerine lessens the capacity of hæmoglobin to take up oxygen, the blood is sometimes chocolate coloured, and yields the spectrum of methæmoglobin.

**Symptoms**—A burning sensation in the throat, nausea, vomiting, giddiness, excessively violent pain in the head, flushing of the face, tumultuous action of the heart, pulsation felt all over the body, prostration, unconsciousness, muscular twitchings, perspiration, stertorous, dyspnoæal breathing, and cyanosis, with complete paralysis, have been observed.

The **fatal dose** is not known, about one ounce caused death in four hours. A man recovered, after extremely violent and dangerous symptoms, who swallowed a tablespoonful of dynamite (a mixture of nitroglycerine with about one third of its weight of siliceous earthy matter), to which a few extra drops of nitroglycerine were added. A man committed suicide by eating two "bobbins" of dynamite four inches long by three quarters of an inch thick.

## CHLORAL HYDRATE.

**Chloral Hydrate** ( $C_2H_3Cl_3O_2$ ) in poisonous doses produces profound coma and abolishes the reflex irritability of the spinal cord. It lowers the blood-pressure partly by paralysing the vasomotor centre and partly by its action on the cardiac ganglia. The absence of chloroform from the expired air of animals to which large doses of chloral hydrate have been given, together with Hammersten's experiments on the blood of animals under the influence of the

<sup>1</sup> *The Lancet*, 1889.

<sup>2</sup> *Brit Med Journ*, 1891.

<sup>3</sup> *Centralbl f klin Med*, 1888.

poison, tend to show that chloroform is not liberated by decomposition of chloral hydrate in the system, as Liebreich, the introducer of this hypnotic into medicine, originally taught. Hammeisten's view is generally accepted, but it appears probable, at any rate exceptionally, that chloral hydrate after being swallowed may liberate chloroform. In a recent fatal case of poisoning with about seventy grains of chloral hydrate, it was observed that the expired air had the odour of chloroform, this is not a solitary experience.

**Symptoms.**—Shortly after the reception of a poisonous dose of chloral hydrate the patient, without any preliminary excitation, becomes drowsy and gradually passes into a state of coma, from which he cannot be roused. Respiration is slow and laboured, long intervals sometimes occurring between the breaths, the pulse is thready and in the later stage slow. The pupils are generally contracted, the face is sunken and cyanotic, or it is pallid and ghastly, the surface of the whole body, but especially of the limbs, is remarkably cold, and is bedewed with perspiration. The reflexes are abolished, and there is absence of sensibility. In fatal cases the temperature is further lowered, and death takes place from heart-paralysis. Cases have been recorded in which the temperature was elevated, Levinstein<sup>1</sup> saw a man, half an hour after he had swallowed 370 grains of chloral hydrate, whose temperature was 103° F. It is characteristic of poisoning by chloral hydrate that the toxic symptoms come on very suddenly, sometimes directly after the poison is swallowed, in these cases the rapidity with which the fatal symptoms appear points to immediate paralysis of the heart, which causes death before the usual effects of the poison have time to manifest themselves. This exceptionally speedy onset of symptoms may possibly be due to rapid splitting-off of chloroform from the chloral.

In cases of impeded circulation through the lungs, or of fatty heart, small doses produce toxic effects. In the case of *Reg v Parton* (Manchester Assizes, 1889), the prisoner was convicted of having caused the death of an elderly man by the administration of chloral hydrate in beer, with the object of robbing him whilst insensible, the man was found in a cab in a state of unconsciousness, and he died shortly after. Post-mortem examination revealed nothing characteristic, the heart was covered and infiltrated with fat, and death probably resulted from heart-paralysis. In the course of the day the deceased had drunk alcohol freely, but there were no indications of death from acute alcoholism. Traces of chloral hydrate were found in the contents of the stomach, the dose taken was probably small but it was sufficient to paralyse a feeble heart.

**Fatal Dose.** The toxic action of chloral hydrate is extremely irregular. Twenty grains caused the death of a patient (who took it for neuralgia) in half an hour, in another case thirty grains were fatal. Kane<sup>2</sup> states that ten grains rendered a woman of thirty-four profoundly comatose, with contracted pupils, she recovered. An old lady of seventy died in nine and a half hours after taking ten grains. Three grains caused the death of a child one year old. On the other hand, recoveries are numerous after enormous doses of several hundred grains, in one instance recovery took place after 420 grains were taken in one dose. In another case, recorded by Acker,<sup>3</sup> a woman swallowed 340 grains of chloral hydrate and the same quantity of potassium bromide in one dose, and recovered. Death has occurred in fifteen minutes, it may be delayed for six or more hours. In a fatal case recorded by Plummer<sup>4</sup> a boy aged sixteen survived nearly forty hours after swallowing more than an ounce of solid chloral hydrate, treatment

<sup>1</sup> *Viertelejahrsschr f ger Med*, 1874

<sup>2</sup> *N Y State Journ of Med*, 1903

<sup>3</sup> *New York Med Rec*, 1880

<sup>4</sup> *The Lancet*, 1894

being resorted to in the interval, the temperature went up to 103° F, and for eighteen hours after the poison was swallowed the patient's breath had a strong odour of chloroform

Chloral hydrate to a great extent is decomposed in the organism, one product—urochloral acid conjugated with glycuronic acid—is found in the urine, in which also small amounts of unchanged chloral hydrate may sometimes be detected

**Treatment.**—The stomach should be emptied by the tube or by an emetic Warmth is of the utmost importance, it should be maintained by hot bottles, and the body should be surrounded by blankets, underneath which friction may be applied Persistent attempts at rousing the patient should be made by means of the faradic current and other usual methods If the breathing fails, artificial respiration should be performed Hypodermic injections of strychnine ( $\frac{1}{2}$  grain) have been recommended, but strychnine is not so good an antidote to chloral hydrate as chloral hydrate is to strychnine Stimulants will probably be required either hypodermically, or alcohol by the mouth or the rectum, hot coffee is useful

**Post-mortem Appearances.** There is no characteristic appearance, the heart and lungs have been found to correspond with the usual conditions of these organs when death has resulted from cardiac or respiratory failure In a few instances, the mucous membrane of the stomach has been found softened, reddened, and easily detached The blood is usually fluid, but not invariably so It has been stated that putrefactive changes are retarded in chloral poisoning, but this effect is not constant

**Chemical Analysis**—The contents of the stomach should be digested for twenty four hours with about three times their volume of absolute alcohol acidulated with a few drops of sulphuric acid, the mixture being repeatedly agitated, the alcoholic extract is then separated, and the alcohol evaporated, the residue is first extracted with petroleum ether, to remove fats (chloral hydrate is insoluble in petroleum ether), and is then shaken out with ethylic ether, which dissolves the chloral hydrate and deposits it on evaporation Urine also may be first treated with petroleum ether and then with ethylic ether in order to extract chloral hydrate As chloral hydrate is decomposed in the living organism, it may escape detection

**Tests.**—The most delicate test for chloral hydrate, as such, is ammonium sulphide A drop added to a weak solution of chloral hydrate produces no immediate change, but in a short time the mixture becomes opalescent and gradually acquires a yellowish or reddish milky appearance, very suggestive of urine overloaded with urates and charged with pigment With very dilute solutions of chloral hydrate a good way of applying this test is to fill a test-tube with the solution and to add a *single drop* of ammonium sulphide (preferably that which is dark coloured), and, after mixing, to gently heat in a Bunsen flame the upper stratum of fluid, which at once darkens in colour and subsequently becomes turbid *Excess of ammonium sulphide interferes with the delicacy of the reaction* A still more delicate way is to place a small quantity of the solution of chloral hydrate in a test-tube and to heat it to the boiling-point, then, by means of a pipette, to add gently (without agitation) one small drop of ammonium sulphide, when in a few seconds the solution becomes turbid By this method a marked reaction is obtained with a solution containing only 0.02 per cent of chloral hydrate Unless chloral hydrate is present in very small amount its presence may be demonstrated by adding a few drops of a solution of potassium hydroxide, which decomposes the chloral into chloroform and potassium formate —





the chloroform is recognised by its odour, and subsequently by the production of phenyl-isocyanide (see under Chloroform), and the potassium formate by boiling it in solution with silver nitrate, which it reduces to the metallic state. Trichloroacetic acid also yields chloroform when treated with alkalis. The  $\beta$ -naphthol test (see under Chloroform) may also be used for chloral hydrate from which chloroform is liberated by the potassium hydroxide used to dissolve the naphthol. When chloral hydrate is present in very small amount, the best plan is to place the organic admixture in a flask, render it alkaline with sodium hydroxide, and then carry out the method described in the chemical analysis of mixtures containing chloroform.

Urochloral acid may be obtained from the urine by evaporating it to one-fourth its volume, acidulating with hydrochloric acid, and shaking out with ether, evaporation of the ether yields needle-shaped crystals arranged in stars, an aqueous solution of these crystals reduces Fehling's solution, and turns the polarised ray to the left. Another reducing substance—conjugated glycuronic acid—is found in the urine after chloral hydrate has been taken.

### CHLOROFORM.

**Chloroform** ( $\text{CHCl}_3$ ).—Occasionally the vapour of chloroform is inhaled for the purpose of committing suicide, and from time to time accidental death occurs to an individual who inhales it in order to produce sleep or to relieve pain. Homicide, due to the inhalation of chloroform-vapour, is all but unknown, a case is recorded by Casper-Liman<sup>1</sup> of a man who thus killed his wife and two children. The medico-legal aspect of the administration of chloroform-vapour as an anæsthetic has been discussed on p. 263, and, in relation to rape, on p. 79.

Attention has recently been directed to the occurrence of death at a more or less remote period after the inhalation of chloroform-vapour, due to fatty changes, especially in the heart. Ambrosius<sup>2</sup> states that a woman, after inhaling about six and a half ounces of chloroform for operative purposes, recovered from the narcosis, but died nearly ninety hours after, at the autopsy, fatty changes were found in the heart. Zeoge v. Manteuffel,<sup>3</sup> in the course of about ten years, saw five cases in which death from secondary syncope, due to fatty degeneration of the heart, occurred in from two to eight days after the inhalation of chloroform for surgical purposes. Fraenkel<sup>4</sup> records four cases, in adults, with autopsies and microscopic examination of the viscera, in which fatty changes of heart, liver, and kidneys were found. Two of these cases were under the influence of chloroform for three hours, the amount inhaled being about three and a half fluid ounces, one died suddenly on the eighteenth day, and the other forty hours after the operation. A third inhaled about seven fluid ounces during four hours, she died on the second day. The fourth was two and a half hours under chloroform, and she died on the fifth day. Martheu<sup>5</sup> states that a woman, aged thirty-four, who whilst under the influence of chloroform for teeth-extraction for forty minutes, inhaled about two and a half ounces, she vomited persistently, and became slightly icteric, the extremities were cold, and she had albuminuria and Cheyne-Stokes breathing. The pulse was very rapid, reaching 140 to 150 on the third day, when she died. At the autopsy, the usual fatty changes in the heart, liver, and kidneys were found. Recent

<sup>1</sup> *Handbuch d. ger. Med.*

<sup>2</sup> *Virchow's Arch.*, 1895.

<sup>3</sup> *Petersb. med. Wochenschr.*, 1895.

<sup>4</sup> *Virchow's Arch.*, 1892.

<sup>5</sup> *Berliner klin. Wochenschr.*, 1896.

cases following operations for appendicitis have been reported by Geoghegan<sup>1</sup> and by Rees<sup>1</sup>. Strassmann<sup>2</sup> states, both on the ground of experience on animals and also from the investigation of fatal cases in human beings, that the remotely fatal results are not dependent on the way in which the chloroform is administered, and, therefore, that the administrator cannot be held responsible. He is of the opinion that ether does not produce similar tissue-degeneration.

It will be noticed that, in all the above cases, the patients were kept under the influence of chloroform for a considerable time, some inhaling large amounts. This probably constitutes the most important factor in the production of the fatty changes, and indicates that in lengthy operations ether is a preferable anæsthetic.

Poisoning with liquid chloroform is not common, and is almost invariably the result of accident or of attempts at suicide, its pungent taste and powerful odour render it unfitted for homicidal purposes, although in one case at least it was strongly suspected to have been administered by the mouth with homicidal intent.

**Symptoms.** The effects produced by chloroform swallowed in the liquid form resemble those caused by its inhalation, together with the local effects of the liquid on the mucous membrane of the stomach and bowels, on which it acts as an irritant, producing symptoms of gastro-enteritis. After a poisonous dose is swallowed, vomiting usually occurs, the vomited matters not always yielding the odour of chloroform, in a short time the patient becomes unconscious and presents the appearance of a person deeply under the influence of chloroform administered by inhalation.—The face is pale and cyanotic, the features are sunken, the pupils are insensitive to light, and are frequently dilated, but they may be of normal size, and the entire surface is cold and bedewed with sweat. Unless the tongue is drawn forward the breathing is stertorous and gradually becomes feebler and slower, the pulse is small and slow, the blood-pressure falling considerably. Death may result from paralysis of the respiratory centres or from heart-paralysis.

If the patient recovers consciousness, he complains of a hot, burning pain in the stomach and bowels, he may have diarrhoea, and the motions may be tinged with blood. The liver may be enlarged and tender, and the skin icteric. Death has occurred from heart-paralysis after the patient has recovered consciousness. A case is recorded by Brasch<sup>3</sup> of a man who drank seventy grammes (a fluid ounce and a half) of chloroform, which produced profound unconsciousness lasting for ten hours, he then came to himself and complained of pain in the region of the liver, which was enlarged, sixty-seven hours after the poison was taken he died from heart-failure. Bridgman<sup>4</sup> records the case of a man aged forty-three who swallowed one fluid ounce of chloroform in order to procure sleep, shortly after, he fell asleep, the narcotic effects lasting about six hours. Three or four hours after the poison was swallowed severe abdominal pains occurred, followed in about an hour by blood-stained motions and by the vomiting of blood-stained matter, at this period the narcosis had passed off, the patient being now perfectly conscious and remaining so till twelve and a half hours after the poison was swallowed, when he suddenly became worse and died.

**Fatal Dose.**—The smallest recorded fatal dose swallowed as a liquid by an adult was about seven fluid drachms, one drachm caused the death of a boy four years old. Recovery has taken place after two ounces in one case,

<sup>1</sup> *Brit. Med. Journ.*, 1912

<sup>2</sup> *Berliner klin. Wch.*, 1898

<sup>3</sup> *Deutsche med. Zeitung*, 1890

<sup>4</sup> *The Lancet*, 1897

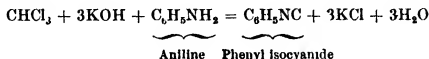
and after three in another (in which the odour of chloroform was present in the breath for two days), and after four ounces in a third case, in all these cases profound narcosis was produced, and in the last general convulsions occurred. Death has taken place in one case one hour after the chloroform (two ounces) was swallowed, in several cases the interval was three hours, from twelve to twenty-four hours is the usual period of survival, and forty-eight hours (after nearly one fluid ounce) is probably the longest.

**Treatment** - When the poison has been swallowed, evacuate the stomach with the tube, and wash it well out. Poisoning by either liquid chloroform or its vapour demands artificial respiration and possibly faradisation of the phrenics, if the breathing fails, administer inhalations of amyl nitrite at frequent intervals, place the patient in the horizontal or even the inverted posture and apply external warmth. To diminish the risk of heart failure the patient should be kept in bed for several hours after recovery of consciousness. Chloroform is principally eliminated by the lungs.

**Post-mortem Appearances.**—Unless the presence of chloroform in the body can be ascertained, there is no characteristic indication of the cause of death. The mucous membrane of the stomach and bowels may be injected, softened and even eroded, incipient fatty changes in the liver, kidneys, and heart have been observed, the blood is frequently fluid, and dark in colour.

**Chemical Analysis -Tests.** The most delicate test is that afforded by splitting up chloroform vapour into chlorine and hydrochloric acid. The substance containing chloroform is placed in a flask furnished with a piece of hard glass tubing, bent at a right angle just above the stopper of the flask, and again twelve or fourteen inches away from the flask, at a right angle, downwards, at a point midway between the two bends a Bunsen flame is allowed to play until the tube is red hot. A second tube, through which air is gently forced, perforates the stopper and dips below the level of the contents of the flask, to which heat is gently applied so as to volatilise the chloroform. When the vapour arrives at the incandescent spot it is split up into chlorine and hydrochloric acid. The former may be recognised by holding to the end of the tube a piece of starch-paper moistened with a solution of potassium iodide, which becomes blue from the action of the liberated iodine on the starch, the latter, by substituting a piece of moist blue litmus-paper, which is reddened, showing the presence of an acid. If the end of the tube is plunged into a solution of silver nitrate, silver chloride is formed, which may be recognised by its insolubility in nitric acid and its solubility in ammonia. The test is so delicate that by finely mincing the lungs, adding sodium carbonate to slight alkalinity, and treating as above described, chloroform has been detected several weeks after death from its inhalation. If present in such an amount that it can be recognised by its odour, chloroform may be separated by distillation from organic admixture, otherwise it is better to adopt the process above described.

After separation chloroform may be tested thus - A little alcoholic solution of potassium hydroxide is put into a test-tube along with ten or twelve drops of aniline, and a little of the chloroform-containing fluid, and is well shaken up. On gently warming the mixture for a short time, the suffocating and disagreeable odour of phenyl-isocyanide or isonitrile is developed. The reaction is thus expressed--



Fill a dry test-tube half an inch deep with  $\beta$ -naphthol and dissolve it in a very small quantity of a strong solution of potassium hydroxide, add some of the suspected fluid—when very dilute the amount added must exceed that of the reagent by one or two volumes— and gently warm in the Bunsen flame. If chloroform is present the liquid turns blue. Chloroform reduces Fehling's solution.

**Quantitative estimation** may be made by introducing the organic admixture into a flask as before described, and passing the vapour through an incandescant combustion-tube containing small fragments of pure caustic lime, with which the chlorine combines. The lime is afterwards dissolved in dilute nitric acid and the chlorine precipitated with silver nitrate—100 parts of silver chloride correspond to 27.758 parts of chloroform.

## BROMOFORM.

**Bromoform** ( $\text{CHBr}_3$ ) closely resembles chloroform in taste and colour. It has a specific gravity of 2.13 and is only slightly soluble in water, two properties which have made it a dangerous remedy on more than one occasion when prescribed for whooping cough. Dean<sup>1</sup> records the case of a girl, aged four, who was taking a mixture containing bromoform, through neglect of shaking the bottle she swallowed from fifteen to twenty drops which had fallen to the bottom, and in half an hour she was unconscious, was livid, and breathed stertorously, the pupils were contracted to pin point size. The stomach was evacuated, and recovery took place. In a fatal case (communicated by Dean) a girl aged five years was given the last dose out of a bottle which when dispensed contained thirty-six minims of bromoform. She became unconscious in twenty minutes and remained so for five hours afterwards, when she died. The fatal dose could not be determined, it was probably from twenty to thirty minims. Dwelle<sup>2</sup> records the case of a child two years old who, after swallowing from thirty to forty drops of bromoform, became unconscious, the heart-beats could not be felt and the breathing was gasping, the breath having the odour of bromoform, the pupils were widely dilated. The patient became cyanosed and, notwithstanding hypodermic injections of strychnine and alcohol, with artificial respiration, death occurred in three to four hours. Kiwull<sup>3</sup> records the death of a child, three years of age, about an hour after the last dose of a mixture containing bromoform had been swallowed. Muller<sup>4</sup> records the case of a child, two years of age, who swallowed six grammes (thirty-three minims) of bromoform. Unconsciousness and convulsions rapidly occurred, respiration was arrested, the pulse was feeble, the pupils were contracted, the surface was cyanotic, and the muscles were relaxed. Death took place in about four hours. On section, the mucous membrane of the stomach and duodenum was found to be injected and oedematous, the odour of bromoform was perceptible when these viscera were opened, and the mucous membrane with which the bromoform had come in contact was injected. The vessels of the brain were injected, and also the meninges, which were of a dark red colour. The blood was thin and liquid. Bommel<sup>5</sup> saw an infant ten months old that recovered (under treatment) after swallowing between fifty and sixty drops of bromoform, in addition to the above named symptoms the muscles of respiration were partially paralysed, and there was trismus, with spasms of the limbs. Czygan<sup>6</sup> states that a boy two and a half years old swallowed between one and two drachms of bromoform, insensibility, slight cyanosis, depressed temperature, contracted pupils, loss of corneal reflex, and extremely feeble respiration occurred, the pulse was over 130 to the minute. With the aid of artificial respiration and of injections of ether and strychnine recovery took place. Berger<sup>7</sup> records two cases, also in children, in which recovery took place with the aid of artificial respiration and hypodermic injections of camphor.

## SULPHONAL.

**Disulphonethyl-dimethylmethane**, or sulphonal, is used to procure sleep. It is a crystalline substance, produced by oxidation of a mixture of ethyl mercaptan and dimethylketone (acetone). It is sparingly soluble in water and ether, and is more freely so in alcohol. As is the case with all hypnotics, sulphonal is used by patients on their own responsibility, and in this way has given rise to serious results.

<sup>1</sup> *The Lancet*, 1893

<sup>2</sup> *Journ. Amer. Med. Assoc.*, 1903

<sup>3</sup> *Centralbl. f. in Med.*, 1902

<sup>4</sup> *Munchener med. Wochenschr.*, 1898.

<sup>5</sup> *Deutsche med. Wochenschr.*, 1896

<sup>6</sup> *Ibid.*

<sup>7</sup> *Munchener med. Wochenschr.*, 1896

In a fatal case recorded by Knaggs,<sup>1</sup> a man took rather more than an ounce of sulphonal. He became comatose, with slow respiration, slow pulse, sometimes increasing to 90 beats per minute, and elevated temperature—ranging from 100° to 103° F., the pupils were of normal size and reacted to light, there was profuse perspiration and total suppression of urine. He remained in the same condition for three days, when the breathing became short and jerky, and finally ceased. Hoppe Seyler and Ritter<sup>2</sup> record the case of a man aged twenty-three, who swallowed fifty grammes (about an ounce and a half) of sulphonal. He slept heavily, the pulse was 120 to 130, and the respirations were 32, the reflexes were abolished. Death from cardiac failure took place seventy hours after the poison was swallowed. After death, reddening of the œsophagus, hyperæmia of the liver, spleen, and kidneys, and ecchymoses in the stomach and duodenum were found. In a case recorded by Reinfuss,<sup>3</sup> a woman, aged forty-seven, took sulphonal in doses of fifteen to twenty two grains almost daily until the total amount reached between two and three ounces. She began to vomit, and complained of pain in the stomach and abdomen, she then lost power over the legs, and had two attacks of clonic spasms, the pupils were contracted and equal, and reacted to light, thirst, diminution in temperature, profuse perspiration, and during the last twenty four hours muscular tremors, and unconsciousness ended in death on the thirteenth day after the commencement of the vomiting. A peculiar appearance observed in the urine, which from the first was dark, and reddish brown in colour, was attributed to the presence of hæmatoporphyrin, albumin and renal epithelium were present. Hanimarsten<sup>4</sup> has shown that the dark colour of urine after drugs such as sulphonal and trional have been taken to excess is not due to the hæmatoporphyrin that is present, but to some other pigment or pigments. When all the hæmatoporphyrin has been extracted, the colour of the urine remains unaltered. Kober<sup>5</sup> relates the case of a man, aged fifty, who for four or five weeks took doses of from seven to twenty two grains of sulphonal, as in the last case, the urine varied in colour from burgundy red to reddish-black, it contained albumin and casts, but no red corpuscles, later on suppression of urine occurred, and the patient died. In addition to hæmatoporphyrin and albumin, Jolles<sup>6</sup> found unchanged sulphonal, and increase of the combined sulphuric acid in the urine. In the case of a man aged thirty-four, who took sulphonal in very large doses—a teaspoonful at once—Ullmann<sup>7</sup> observed reeling gait, stammering speech, and erratic gesticulations, which lasted four or five days. In another case there were ataxic symptoms which prevented locomotion and impeded speech. Treilian<sup>8</sup> saw a woman aged twenty-eight, who became ataxic and cyanosed, with very slow, shallow breathing, and feeble and intermittent action of the heart, vomiting occurring several times, after taking twenty grains of sulphonal one night and fifteen grains the following night. The patient recovered. In a case recorded by Whatley,<sup>9</sup> a man fifty years of age suffered from circular patches of erythema over various joints, six or seven hours after taking a single twenty grain dose of sulphonal, thirty hours after, serum exuded as from a small blister.

On the other hand, enormous doses have been taken without any permanent ill effects. Neisser<sup>10</sup> records a case in which a youth, aged fifteen, for the purpose of committing suicide, took fifty grammes of powdered sulphonal, and shortly after another fifty grammes, amounting together to more than three ounces. In three quarters of an hour he became unconscious, and was found about six hours after, when he was taken to the hospital, emetics were administered, and the stomach was washed out, he slept until the sixth day, and on the ninth day was perfectly well. The urine contained neither albumin nor sugar, but unchanged sulphonal was obtained from it. In another case a man, after taking over an ounce, slept for five days and then recovered.

**Tests.**—If a little dry sulphonal is heated in a test tube with charcoal or iron in powder, the odour of mercaptan is given off, if iron is used, the subsequent addition of hydrochloric acid to the residue liberates sulphuretted hydrogen. When a little dry sulphonal is melted and the heat is continued until the clear liquid boils, the addition of pyrogallol produces a brown colour with evolution of mercaptan.

## TRIONAL.

**Diethylsulphonmethylethylmethane**, or trional, has been recommended as a substitute for sulphonal as being less liable to produce ill effects. Like all such synthetically formed drugs, its use is not free from danger, at least six cases of alleged poisoning have been recorded.

<sup>1</sup> *Brit Med Journ*, 1890.

<sup>2</sup> *Munchener med Wochenschr*, 1897.

<sup>3</sup> *Wiener med Blätter*, 1892.

<sup>4</sup> *Skand Arch f Physiol*, 1891.

<sup>5</sup> *Centralbl f klin Med*, 1892.

<sup>6</sup> *Internat klin Rundschau*, 1891.

<sup>7</sup> *Corresp Blatt. f Schweiz Aerzte*, 1889.

<sup>8</sup> *Brit Med Journ*, 1899.

<sup>9</sup> *The Lancet*, 1904.

<sup>10</sup> *Deutsche med Wochenschr*, 1891.

Repeated doses have caused sore throat, coryza, giddiness, uncertainty of gait, and dementia. Schulze<sup>1</sup> states that a woman, aged fifty four, took a series of doses of from seven to twenty three grains of trional daily for about a month—in all about 380 grains. She began with anorexia, followed by pain in the abdomen and vomiting, the urine contained hæmatoporphyrin, death occurred a few days after the last dose. Herting<sup>2</sup> records the case of a woman, aged thirty six, who took 340 grains in a series of doses lasting over twenty-four days, she had previously been taking sulphonal. The symptoms were as in the preceding case, the urine becoming dark in colour from the presence of hæmatoporphyrin, the patient died twenty six days after the first appearance of the dark coloured urine, and fifteen days after the last dose of trional. Berger<sup>3</sup> relates a case in which sixty grains of trional taken in the twenty four hours produced dangerous symptoms in a medical man—extreme somnolence, stertorous breathing, which occasionally became of the Cheyne Stokes type, cyanosis, hallucinations, and dark coloured urine. Collatz<sup>4</sup> states that 120 grains of trional, taken in a single dose with suicidal intent, were recovered from, there was no hæmatoporphyrin in the urine. Mackintosh<sup>5</sup> reports a recovery after 18 grains had been taken, together with 20 grains of veronal.

### VERONAL.

**Diethyl malonyl urea**  $[(C_2H_5)_2C(COHN)_2CO]$  Malourea or veronal has been responsible for a considerable number of fatal cases of poisoning in recent years. Knowledge of the drug and its effects has recently been summarised by Willcox.<sup>6</sup> Veronal is a white crystalline powder, melting at  $191^\circ C$ , and possessing a bitter and somewhat nauseous taste, and an acid reaction to litmus. It is only slightly soluble in cold water, but at  $100^\circ C$  1 part is soluble in 15 of water, it is soluble in alcohol, ether, acetone and alkaline solutions.

Veronal is a powerful hypnotic, and owing to belief in its comparative freedom from danger there is no doubt that it has been frequently taken for insomnia without advice or prescription from a medical man. It is now in Part II of the Poisons Schedule. The pharmaceutical dose is 5 to 10 grains, and this should not be exceeded except under very special circumstances.

**Symptoms.**—Willcox gives the following description. After a single large dose, headache, drowsiness, sometimes with ataxy and reeling gait, occur, the patient falls into a deep sleep, from which he can only be aroused with difficulty. In severe cases, the sleep deepens into coma with cyanosis and rapid, often stertorous, breathing. In this stage, it is very common for a marked rise of temperature to  $103^\circ F$  or even more to occur, and physical examination of the lungs will probably show areas of dulness on percussion, increased bronchial breathing and moist sounds, accompanied or followed by the signs of general œdema of the lungs such as moist bubbling râles all over. On several occasions this condition has led to veronal poisoning being mistaken for pneumonia. Recovery when these conditions develop is uncommon, sometimes the coma and high temperature will last for as long as four days before death supervenes, but after a very large dose death may occur in less than twenty-four hours. Skin rashes, erythematous in type, or of a rubefoliform or scarlatiniform character, have been described, and it is stated that urticarial rashes and also extreme pruritus and œdema of the face occur. But affections of the skin are not common. Urinary disturbances such as suppression albuminuria, hæmatoporphyrinuria and hæmaturia have also been described, but are not common.

In chronic veronal poisoning (veronal habit), the patient often develops an abnormal mental condition, the whole mental balance may be upset, visual

<sup>1</sup> *Deutsche med. Wochenschr.*, 1894.

<sup>2</sup> *Ibid.*, 1894.

<sup>3</sup> *Munchener med. Wochenschr.*, 1895.

<sup>4</sup> *Berliner klin. Wochenschr.*, 1893.

<sup>5</sup> *Lancet*, 1910.

<sup>6</sup> *Internal Med. Congress*, London, 1913.

hallucinations are common, and delusion may occur with complete disorganisation of the moral sense, as in the morphia and cocaine habits. Tremors and marked ataxy are common, and the condition may resemble cerebellar disease. The speech is often thick and indistinct, and disturbances of sight may occur. The gait may resemble that of a drunken person. Veronal is more toxic in cases of renal disease than in normal persons, and it is important that constipation should be avoided if the drug is being given frequently, as otherwise toxic symptoms may develop.

**Fatal Dose.**—Death has been recorded after doses so small as ten or fifteen grains, but in these cases other factors were probably present. It may be taken that a dose of fifty grains is dangerous in a healthy adult, and might be regarded as the average minimum lethal dose.

**Treatment.**—If the patient is seen within four hours of taking the drug, the stomach should be well washed out with warm water. After the last washing, a pint of hot strong coffee with some milk in it and an ounce of castor oil should be introduced into and left in the stomach. Cardiac stimulants, such as strychnine hydrochloride gr.  $\frac{1}{10}$  with digitalin gr.  $\frac{1}{100}$ , may be given every four hours. Warm normal saline should be given subcutaneously, and also rectal injections of normal saline containing 4 per cent. of glucose, in amounts of fifteen ounces every four hours. If there is much cyanosis, oxygen should be given, and when the pulse is feeble the oxygen may with advantage be passed through a wash-bottle containing absolute alcohol since the combination of oxygen and alcohol vapour is a valuable cardiac stimulant. During the comatose state it may be necessary to draw off the urine with a catheter.

**Post-mortem Appearances.**—These are not characteristic. Cyanosis is often present and post-mortem staining well marked. The heart usually shows marked dilatation, the right side being more affected than the left. The lungs show marked hypostatic congestion, and patches of pneumonic consolidation are frequently present. The brain and abdominal viscera are usually congested.

**Tests.**—These are not very satisfactory. The melting point of  $191^{\circ}\text{C}$  may be employed, but the drug must be isolated in a pure condition. No precipitate is produced by the reagents ordinarily used to precipitate alkaloids. A solution of veronal if boiled with a 20 per cent. solution of caustic potash is not decomposed, and no brown colour is produced by the addition of Nessler's reagent. If solid veronal is fused with caustic potash a disagreeable rancid smell is caused, and on adding water and then Nessler's reagent a well-marked brown colour is produced. Two drops of dilute nitric acid and then Millon's reagent added to a solution of veronal give a white gelatinous precipitate soluble in excess of the reagent. Veronal may be extruded from the tissues by treatment with 95 per cent. alcohol just acidified with acetic acid.

## CARBON BISULPHIDE.

**Carbon Bisulphide** ( $\text{CS}_2$ ) has only exceptionally caused **acute** poisoning, **chronic** poisoning, by its vapour, is more common in consequence of its widespread use in india-rubber and gutta-percha works.

**Symptoms of Acute Poisoning.**—In a case recorded by Davidson<sup>1</sup> a man swallowed about two ounces of carbon bisulphide. When seen, the patient was in a state of collapse, the muscles were relaxed and the pupils dilated and insensible to light, the pulse was quick and feeble, the breathing laboured,

<sup>1</sup> *Med Times and Gaz*, 1878

the odour of the poison being perceptible in the breath the lips were blue, and the surface was cold, occasional convulsive tremors or shiverings occurred, the odour of carbon bisulphide could be perceived both in the urine and the fæces. Recovery took place, the patient being well on the fifth day. A fatal case is recorded by Foreman<sup>1</sup>. A man swallowed about half an ounce of carbon bisulphide, and became comatose in half an hour. The respirations were slow and laboured, the pulse 150 to 160 in the minute, the surface was cold and clammy, the pupils were normal, death occurred two and a quarter hours after the poison was swallowed. At the necropsy, the odour of the poison was perceptible, the posterior wall of the stomach was congested for about the size of a crown-piece, and hæmorrhagic points were visible in the gastric mucous membrane, the veins were gorged with black blood, the blood throughout being fluid. The urine yielded the odour of carbon bisulphide.

**Treatment.**—The stomach-tube should be used, and the drowsiness then combated as is customary in poisoning with other narcotics. Warmth should be applied to the body, and stimulants administered internally, and in case of need, artificial respiration should be resorted to. If there is no diarrhœa, purgatives should be given.

Carbon bisulphide is eliminated by the lungs, kidneys, and bowels. Death is apparently due to paralysis of the respiratory centres, which is also the cause of death in animals experimentally poisoned with carbon bisulphide.

**Chemical Analysis.**—From organic admixture carbon bisulphide may be separated by distillation. It is recognised by its odour, and by giving a black precipitate of lead sulphide when heated with lead acetate and potash.

**Chronic Poisoning.**—Like many other volatile poisons, carbon bisulphide, when repeatedly inhaled, produces peripheral neuritis, in addition to disorders of the digestive tract. Wiener's<sup>2</sup> experiments show that a mean of 23·7 per cent. of the inspired vapour of carbon bisulphide is absorbed. In the early stage of chronic poisoning as it occurs amongst workmen in certain departments in rubber factories, the appetite fails, and the patient is always conscious of the odour of the bisulphide, even when away from his work, sometimes a state of mental exaltation or of depression occurs, with sleeplessness, headache, nausea, vomiting, and colicky pains. These symptoms are followed by those of neuritis, in some cases investigated by Ross,<sup>3</sup> the earliest of the neural symptoms was a burning sensation in the hands, alternating with numbness, then followed tingling and numbness in the feet, with weakness. In one case the immediate narcotic effects of the vapour were demonstrated by the patient's desire to return to his work, because when he inhaled the vapour he felt some relief from the symptoms. The extensor muscles of the forearm and those of the leg were atrophied and partially paralysed, producing wrist-drop and ankle-drop respectively, and the patient experienced numbness and tingling in the toes and feet, and in the tips of the fingers, the field of vision was restricted for all colours. Another patient was disturbed by horrible dreams, fancying himself surrounded by animals, sometimes, when at his work, he found himself talking nonsense. When the vapour of carbon bisulphide is present in considerable amount, the workmen not unfrequently develop symptoms resembling those of delirium tremens. Amblyopia and scotomata without retinal changes are common, and there may be considerable diminution in the number of the red blood corpuscles. Krudener<sup>4</sup> reports commencing optic atrophy in a man who worked with carbon bisulphide in a chemical factory.

<sup>1</sup> *The Lancet*, 1886.

<sup>2</sup> *Dissert.*, Würzburg, 1906.

<sup>3</sup> *Med Chron.*, 1887.

<sup>4</sup> *Zeitschr. f. Augenheilk.*, 1906.



## PETROLEUM AND PARAFFIN OIL.

**Petroleum** is a natural oil, consisting of a mixture of the higher, but not the highest, paraffins or hydrocarbons of the series  $C_nH_{2n+2}$ , of which marsh-gas is a type. The commercial oil varies in sp gr from 7 to 825, and in boiling point from  $150^{\circ}$  to  $300^{\circ}$  C. If well refined it is transparent, slightly fluorescent liquid, free from colour, but it usually presents a slight yellowish tinge when examined in bulk, it has a peculiar penetrating odour. It is largely used as an illuminant. An early product of distillation of the crude oil is known as petroleum ether, which has a sp gr of 66 to 67, and a much lower boiling-point ( $50^{\circ}$  to  $60^{\circ}$  C) than the ordinary variety. Petroleum ether is used for extracting fats from organic fluids and as a solvent.

**Paraffin Oil** is a mixture of paraffins obtained by the distillation of shale, from the toxicological standpoint it does not differ from petroleum. In retail trade petroleum and paraffin oil are sold indifferently, the one for the other.

The toxic properties of petroleum depend greatly upon the kind of oil taken, but in any case it is not an active poison. Lewin,<sup>1</sup> basing his conclusions on a number of experiments on animals, and observations on the human subject, does not regard petroleum as a poison in the ordinary sense of the word, the dose necessary to produce toxic effects being so large. Pure paraffin is probably quite innocuous. McCulloch<sup>2</sup> saw a man, aged forty-three, one hour after he had swallowed nearly half a pint of paraffin oil, he was pale, and his breath had the odour of the oil, he complained of his throat being hot and dry, and he had a feeling of warmth in the epigastric region, but no pain, the pupils were normal, the pulse was full, but this probably resulted from excitement. After an emetic he vomited a little food and about eight ounces of paraffin oil. The oil was in his stomach an hour, along with very little food, but it did not cause the least indication of gastric irritation, the man was quite well the next morning. On more than one occasion a pint of petroleum has been swallowed without causing more than temporary disorder. On the other hand, Vincent<sup>3</sup> saw a girl, aged fifteen and a half, who, fifteen or twenty minutes after swallowing about half a pint of paraffin oil, vomited, and was cold, with pale, anxious face, feeble pulse—132 in the minute—and sighing respiration, she had pain in the throat, epigastrium, and left hypochondrium, recovery took place. Unusually severe symptoms occurred in the case of a woman, aged thirty-six, who was seen by Carruthers.<sup>4</sup> During a debauch she swallowed half a cupful of paraffin oil, in half an hour she had violent pain, and vomited, the vomit was stated to contain blood. When seen three or four hours later, she complained of pain in the epigastrium and in the left lumbar region, although she had vomited in the interval, the matter subsequently vomited proved that a considerable quantity of oil had been retained, its odour was present in the breath for twenty-four hours. The motions contained blood, along with paraffin oil, and a considerable quantity of oil floated on the surface of the urine after it had stood a while, on distillation 6 c.c. of pure oil were obtained, subsequently the urine contained albumin and blood. The patient was well in a week. Some writers doubt the possibility of petroleum being present in bulk in the urine, Lewin, in his experiments on animals, found that it did not appear as such in the urine, and disputes its occurrence in the human subject. Several cases are recorded, however, besides the one just related,

<sup>1</sup> Virchow's Arch., 1888.

<sup>2</sup> The Lancet, 1885.

<sup>3</sup> Brit. Med. Journ., 1868.

<sup>4</sup> The Lancet, 1890.

in which, after the individual had swallowed petroleum in large amount, unaltered oil was found floating on the urine

Johannsen<sup>1</sup> records a fatal case in which a girl aged twelve drank an unknown quantity of American petroleum oil, she became cyanotic with laboured respirations, 50 to the minute, the pulse being 144. She vomited, and both the vomit and the motions contained petroleum, the urine was lost, the patient became drowsy, and in five or six hours died in an unconscious state. On section the stomach was found distended with air, when opened it yielded a strong odour of petroleum, which was perceptible along the intestines as far as the jejunum, the gastro-enteric mucous membrane was pale without any sign of injection or of excoriation. M'Dougall<sup>2</sup> saw an infant aged fourteen months that died one hour and fifty minutes after swallowing one ounce and a quarter of paraffin oil, in this case convulsions and cyanosis occurred, but no vomiting. Lesser<sup>3</sup> examined the body of an infant, a year and a half old, in which death, from extreme œdema of the glottis, occurred five hours after swallowing a small quantity of petroleum. Death may ensue from secondary effects, as in the case recorded by Lugol<sup>4</sup>. A woman aged forty swallowed a "glassful" of petroleum, the pulse became small, there was neither nausea nor vomiting, but some discomfort was felt in the stomach. On the surface of a motion passed a few hours after, some petroleum floated, which readily took fire on the application of a light. Gastro-enteritis set in, and the patient died on the twentieth day. Biller<sup>5</sup> records the case of an infant, eighteen months old, that drank some "gasoline", it became unconscious, powerless, livid in the face, tympanic in the abdomen, and cold on the surface, death occurred in thirty minutes.

**Vaseline**, a mixed intermediate product between the liquid and solid members of the paraffin series of hydrocarbons, is usually regarded as a harmless substance, in one instance, however, recorded by Robinson,<sup>6</sup> half a teaspoonful given to each of three children for sore throat produced vomiting, pains in the knees, and cramps in the legs, with partial collapse, all the children recovered.

### TETRACHLORETHANE, $C_2H_2Cl_4$

This substance was one of the constituents of a "dope" or varnish which was largely used in aeroplane factories during the early years of the war for covering the wings of aeroplanes. As its toxic nature had not been recognised, no special precautions were at first observed in its use. In November, 1914, however, a series of 19 cases of jaundice, one of which was fatal, in an aeroplane factory, led to further investigations, and the condition was made notifiable. In all, some 70 cases with 12 deaths were reported. The symptoms were loss of appetite, nausea, vomiting, headache, drowsiness, and jaundice, with delirium in some cases. Post-mortem, there were degenerative changes in the liver, and marked reduction in its size. The establishment of exhaust ventilation by means of fans appreciably improved the conditions, but isolated cases continued to occur until July, 1916, when a substitute for tetrachlorethane was found, and dope containing this ingredient ceased to be used.

<sup>1</sup> *Berliner klin. Wochenschr.*, 1896.

<sup>2</sup> *Med. Chron.*, 1898.

<sup>3</sup> *Vierteljahrschr. f. ger. Med.*, 1898.

<sup>4</sup> *Repertoire de Pharmacie*, 1871.

<sup>5</sup> *New York Med. Journ.*, 1889.

<sup>6</sup> *Brit. Med. Journ.*, 1886.

## CHAPTER XXXIV

## CARBON COMPOUNDS—AROMATIC GROUP.

## BENZENE AND ITS DERIVATIVES.

**Benzene** ( $C_6H_6$ ), or benzole, is one of the principal ingredients in coal-tar, from which it is obtained by fractional distillation, commercial benzene contains small quantities of some of the other light hydrocarbons. It is a colourless, volatile liquid, having an odour which recalls that of coal-gas, it is exceedingly inflammable, and gives off a vapour which is explosive when mixed with air, it is insoluble in water, on which it floats. It is used in the manufacture of aniline, and also in cleaning gloves and wearing apparel.

**Symptoms of acute poisoning** much resemble those of alcohol—a stage of excitement, which is quickly followed by heaviness in the head and a tendency to stupor or coma. Foulerton<sup>1</sup> records the case of a man who, in the pursuit of his avocation, entered a large tank in which benzene-vapour had accumulated, when found, he was insensible, and could not stand, he could answer questions in an indistinct way, moaning and laughing hysterically. His face was flushed, and the surface of the body was cold, there were muscular twitchings, the pupils were dilated, and reacted to light, the pulse was 88, full and soft, the respirations, reduced to 8 or 9 in the minute, were deep, stertorous, and irregular, as much as fifteen seconds sometimes intervening between the breaths. The patient vomited, the ejected matter smelling of benzene. Recovery took place. A fatal case of poisoning by benzene-vapour is recorded by Sury-Bienz<sup>2</sup>. It occurred to a workman in a chemical manufactory, who had to attend to a process in the course of which a great quantity of benzene was volatilised. He was heard to call out that he was on fire, he then reeled, fell to the ground, and died forthwith, a fellow workman who ran to his help perceived a powerful odour of benzene, but there was no fire. At the necropsy the veins were found filled with fluid blood, and there was some œdema of the lungs, but nothing else of moment.

Averill<sup>3</sup> met with a case, in which a man accidentally swallowed three or four drachms of benzene, he became pale and unconscious, with small, weak, rapid pulse, and slow breathing, the pupils did not react to light. The stomach was evacuated, and the vomited matter contained oily-looking globules, which took fire on applying a light. Recovery occurred, the odour of benzene being perceptible in the breath sixty-two hours after the poison was swallowed. Falk<sup>4</sup> records a fatal case in a child two years old, death taking place ten minutes after a mouthful of benzene was swallowed, the post-mortem appearances, beyond a faint odour of benzene on opening the abdomen, were negative. Kelynack<sup>5</sup> records a fatal case which occurred in a woman aged twenty-six who drank about an ounce of benzene. When seen six hours after, she was unconscious,

<sup>1</sup> *The Lancet*, 1886<sup>2</sup> *Brit Med Journ*, 1889<sup>3</sup> *Vierteljahrsschr. f. ger Med*, 1888<sup>4</sup> *Vierteljahrsschr. f. ger Med*, 1892<sup>5</sup> *Med Chron.*, 1893.

the pulse was very quick and feeble, and the respirations were rapid. The pupils were slightly contracted and reactionless, the extremities were cold, and the lips, ears, and nose distinctly blue. On washing out the stomach the liquid which returned had a strong odour of benzene. After subcutaneous injections of ether and strychnine the patient became sufficiently conscious to complain of pain in the abdomen and severe frontal headache, she also suffered from nausea, diarrhoea then set in, and death from cardiac failure took place twelve hours after the poison was taken. At the necropsy the various cavities and organs yielded a strong odour resembling that of aniline, on slitting up the intestines an odour like that of coal-gas was perceived, the tissues generally were hyperæmic, and there were a few hæmorrhages in the bronchi and in the small intestines, but there was no erosion of the alimentary tract. The blood yielded the spectrum of oxyhæmoglobin. No trace of aniline could be obtained from the urine. In a case related by Spurr,<sup>1</sup> a woman aged twenty-six drank one ounce of benzene, which produced severe gastro intestinal irritation, with elevation of temperature, hurried respiration and quickened pulse, death took place fifty hours after the poison was swallowed. At the post-mortem examination the œsophageal and gastric mucous membrane was inflamed, and also that of the duodenum for three inches below the stomach, erosions were present. The lungs were congested and the bronchi contained puriform matter. Selling<sup>2</sup> records two fatal cases of purpura among girls employed in a factory where benzol was being used. Hæmorrhages occurred into the skin and mucous membranes. Purpuric spots were exhibited by four other persons working in the same room.

### NITRO-DERIVATIVES OF BENZENE.

**Nitrobenzene** ( $C_6H_5NO_2$ ), or nitrobenzole, is the product yielded by the action of nitric acid on benzene. It is a light yellow liquid having an odour resembling that of oil of bitter almonds, and is known in commerce as **artificial oil of bitter almonds** or **oil of mirbane**, it is largely used in the preparation of aniline, in the manufacture of furniture polish and boot polish, and to scent toilet soaps.

**Dinitrobenzene** [ $C_6H_4(NO_2)_2$ ].—The compound used commercially is the meta dinitro benzene, which, when pure, forms long rhombic prisms of a light yellow tint, the commercial variety is yellowish brown. It is soluble in alcohol and ether and, when impure, to a slight extent in water. It is produced in aniline works and enters into the composition of the explosive "roburite" now largely used in coal mines for the purpose of blasting. **Roburite** consists of a mixture of dinitrobenzene, or chloro dinitrobenzene and ammonium nitrate.

#### Nitrobenzene.

**Symptoms** of acute poisoning with **nitrobenzene**. The most characteristic symptom is the occurrence of a livid or cyanotic appearance of the face, the lips especially acquiring a dull red colour, the fingers and toes, and even the whole body, may be intensely blue. Numbness of the tongue and salivation may occur. There is giddiness and pain in the head, and if the patient is able to walk, his gait is unsteady and his muscular power feeble, vomiting may occur, the vomited matters probably having the odour of the poison, which also pervades the breath, drowsiness develops and rapidly passes into stupor or coma. Trismus, tetanus, increased knee jerk, and ankle clonus have been observed. The pulse is usually feeble and quickened, and may be intermittent, the breathing is often shallow and irregular, with quickened expiration, the temperature is reduced and the surface is clammy. The pupils have been observed to be fixed—contracted in some cases and dilated in others, dilatation is common in the comatose stage. Blood withdrawn during life has been found darker than usual. In some cases jaundice has occurred on the third or fourth day.

<sup>1</sup> *The Lancet*, 1899

<sup>2</sup> *Johns Hopkins Hospital Bulletin*, 1910

Filehne<sup>1</sup> found that in dogs poisoned with nitrobenzene the blood was chocolate brown in colour, on spectroscopic examination it yielded a band in the red near the hæmatin band, which he regards as directly due to the action of nitrobenzene, in no case did he find that the nitrobenzene was converted in the system into aniline. Filehne explains the dyspnoea as being due to the incapacity of the hæmoglobin to carry oxygen to the tissues, animals poisoned with nitrobenzene exhale more CO<sub>2</sub> and take up less O than in the normal. Lewin<sup>2</sup> found the band in the red to be identical with that of hæmatin.

**Fatal Dose**—Twenty drops have proved fatal. Recovery has occurred after very nearly an ounce, but the patient received prompt and efficient treatment. Wermuth<sup>3</sup> states that a woman swallowed ten drops in order to procure abortion. In sixteen hours the urine was dark coloured, it was lævo rotatory and it reduced Fehling's solution. The blood showed the spectrum of methæmoglobin. Dodd<sup>4</sup> records the case of a man aged forty-seven who swallowed two drachms of nitrobenzene and then ate his dinner, after which he walked three quarters of a mile. One hour and a half after the poison was swallowed extreme cyanosis developed, the skin was clammy, the pulse feeble, and the respirations were shallow, irregular, and sighing, among other symptoms was trismus. The breath had the odour of nitrobenzene. The stomach was well washed out, and the patient recovered. Grant<sup>5</sup> records the case of a woman of forty who took less than half an ounce of oil of mirbane. Coma, marked cyanosis, almost imperceptible pulse, and stertorous breathing followed. Death took place in an hour and a quarter. In fatal cases death takes place in from one to twenty-four hours.

**Treatment**—Evacuate and thoroughly wash out the stomach, apply external warmth and friction, and if necessary use artificial respiration and faradism. Stimulants may be required, they should not, however, be given by the mouth unless the stomach has been thoroughly washed out, as alcohol is a solvent of nitrobenzene. In severe cases transfusion of defibrinated human blood may be resorted to after removal of an equal amount of blood from the patient.

### Dinitrobenzene.

Poisoning with dinitrobenzene usually occurs in manufactories where this substance is used, it is received into the system either in the form of vapour or of finely divided particles floating in the air, or from handling it in bulk. It is customary in many works to provide the men with rubber gloves, as otherwise their hands become contaminated with the poison, which is thus transferred to food, it is probable that prolonged handling of dinitrobenzene may lead to its introduction through the skin.

The symptoms of acute poisoning resemble those produced by nitrobenzene.—Headache, giddiness, loss of power in the limbs, blue coloration of the lips, cold and livid surface, quickened feeble pulse, dyspnoea, shallow irregular respiration, with long intervals between the breaths, and coma. The cyanotic appearance may be limited to the face or it may spread to the limbs, usually the trunk is not much affected, the blood is dark, and sometimes chocolate coloured. Vomiting frequently occurs spontaneously.

**Chronic poisoning** with dinitrobenzene, which also occurs amongst those who prepare or purify it, produces a different class of symptoms. Schroder and Strassmann<sup>6</sup> investigated several cases, and describe the chief symptoms as follows.—Headache, pain in the stomach with irregular action of the bowels, loss of appetite, sleeplessness and general feeling of lassitude. The lips are blue and the skin acquires a dirty yellow tint, the sclera being yellow, in some cases the mucous membrane of the mouth, especially of the uvula and pharynx, looks as though it was covered with a yellow bloom, which, however, cannot be wiped off. The gastric and hepatic regions are very tender on pressure, the liver usually being enlarged. The urine is dark brown, but quite clear, dinitrobenzene has been demonstrated in it. The symptoms present very much the appearance of those due to catarrhal jaundice, but the urine yields no trace of bile, the motions retain their colour, and the prostration and blue colour of the lips are different from anything observed in cases of jaundice. Rohl<sup>7</sup> describes certain effects produced on the nervous system by chronic dinitrobenzene poisoning resembling those of peripheral neuritis.—Numbness, a sensation of cold in the feet, with various paresthesiæ and cramps.

From experimental researches by Huber<sup>8</sup> it appears that dinitrobenzene combines

<sup>1</sup> *Arch f exp Pathol*, 1878

<sup>2</sup> *Virchow's Arch*, 1879

<sup>3</sup> *Biochemisches Centralbl*, 1907

<sup>4</sup> *Brit Med Journ*, 1891

<sup>5</sup> *Brit Med Journ*, 1913

<sup>6</sup> *Vierteljahrsschr f ger Med* (Supp.), 1891

<sup>7</sup> *Ueber acute u. chron Intox durch*

*Nitrokörp d Benzolreihe*, 1890

<sup>8</sup> *Virchow's Arch*, 1891

with hæmoglobin, which then yields a spectrum identical with that of acid hæmatin, but it does not respond like hæmatin to reducing agents. On the addition of ammonium sulphide the band in the red, between C and D, persists but it is slightly displaced, the other two bands remain unchanged. The spectrum is probably the same as that described by Fiehlne in his investigations on nitrobenzene, it is not always obtained with the blood of animals poisoned with dinitrobenzene, nor has it been observed in the human subject. As the result of further researches Strassmann and Strecker<sup>1</sup> state that the length of time during which dinitrobenzene is administered to animals determines the amount of change undergone by the blood, it is most pronounced in very chronic poisoning, and they agree with Huber that destruction of the red blood corpuscles (with consequent hæmoglobinuria and fatty changes in the viscera) is a constant result of chronic dinitrobenzene poisoning.

The urine, as before stated, contains dinitrobenzene, which exists as such, no derivatives have so far been detected. By means of treatment with zinc and hydrochloric acid, the dinitrobenzene present in the urine may be converted into phenylenediamine, if the urine is then made alkaline with soda and shaken out with ether, and the residue after evaporation of the ether is treated with sodium nitrite and acetic acid, a brown colour—Bismarck brown—is produced. If the urine is shaken out with ether without being previously treated with zinc and HCl, no effect is produced on the ethereal residue by sodium nitrite, showing that the dinitrobenzene is not decomposed in the system into phenylenediamine nor into nitraniline.

**Roburite**, being largely composed of dinitrobenzene, produces similar toxic symptoms. Spurgin<sup>2</sup> reports an interesting case of **acute poisoning** with this explosive—A boy of sixteen slept in a room in which roburite had been sprinkled to poison cockroaches, he was found deeply cyanosed—the lips, tongue, and fingers being nearly black, the surface was cold, there was dyspnoea with laboured breathing, and the pulse, 135 in the minute, was very weak. Another lad who slept in the same room was only slightly cyanosed. Recovery took place in both cases.

**Chronic poisoning** with roburite is identical with that due to dinitrobenzene, Ross<sup>3</sup> investigated some cases and found well marked symptoms of peripheral neuritis in addition to the gastro hepatic symptoms.

It is to be observed that the symptoms produced by roburite are due to dinitrobenzene being introduced into the system as such, and not to the fumes produced by detonation of the explosive. See section on the gases given off by explosives.

**Treatment**—In the acute form the treatment is like that for poisoning with mononitrobenzene, excepting that the stomach tube will not be needed unless the poison has been swallowed in bulk. **Chronic poisoning** demands withdrawal from the influence of the poison and general treatment of the symptoms as they present themselves, unless the condition is far advanced, improvement, if not ultimate recovery, takes place.

**Post-mortem Appearances**—When death occurs from acute poisoning with either mono or dinitrobenzene, with the exception of the odour of the poison, the indications are far from being characteristic. The blood has been found dark, or chocolate coloured, ecchymoses have been observed on the mucous membranes, and the internal organs have been found paler in colour than usual, the cyanotic hue of the skin and mucous membrane is not always visible after death. Letheby,<sup>4</sup> who was one of the first to investigate poisoning with nitrobenzene, found the liver purple in colour and the heart and veins as in death from asphyxia.

**Chemical Analysis—Tests**—Nitrobenzene may be separated from organic admixture by distillation, it is distinguishable by its odour. If the distillate is treated with zinc and hydrochloric acid, the nascent hydrogen liberated converts the nitrobenzene into aniline, the product, diluted with water, filtered if necessary and treated with bleaching powder added a little at a time, yields a purple colour which tends to go back. If, after conversion into aniline, potassium hydroxide and a few drops of chloroform are added to the distillate, and it is heated, the presence of phenyl isocyanide may be recognised by its odour. (See tests for chloroform.) If dinitrobenzene was the poison, it may sometimes be detected in the blood by directly converting it into metaphenylenediamine (as described in the case of the urine), shaking out with ether, and after evaporation of the ether testing the residue with sodium nitrite.

**Dinitrotoluene** [ $C_6H_4(NO_2)_2CH_3$ ] produces effects in every way similar to the nitrobenzene compounds, with which it is closely allied. Maceray<sup>5</sup> records the case of a child three years old, who swallowed a piece of dinitrotoluene the size of a pea. One hour after

<sup>1</sup> Friedreich's *Blatter f. ger. Med.*, 1896

<sup>2</sup> *Brit. Med. Journ.*, 1891

<sup>3</sup> *Med. Chron.*, 1889

<sup>4</sup> *Proc. Royal Society*, 1863

<sup>5</sup> *The Lancet*, 1888

there was complete relaxation of the muscles and coma, the breathing was rapid and shallow, the pulse rapid, and the surface cold and cyanotic, the pupils were equal and insensitive to light, convulsions occurred. The cyanosis, which was totally different to that seen in asphyxia, disappeared within twenty-four hours, the lips being the last to regain their natural colour, and the child was quite well the next day.

### TRINITROTOLUENE.

Poisoning by trinitrotoluene [ $C_6H_2(NO_2)_3CH_3$ ] has been the subject of an exhaustive monograph<sup>1</sup> issued by the Medical Research Council, from which most of the following is taken —

This high explosive was largely used during the war for filling shells and mines. Mixed with 40 per cent of ammonium nitrate, it was known as amatol. Although trinitrotoluene had been used in the manufacture of explosives before the war, it had been looked upon as innocuous, and it was not until the end of 1915 that it was recognised to possess toxic properties. The first fatal cases undoubtedly due to T N T was that of a man who worked at shell-filling from May to July, 1915, and died in August of the same year. He developed jaundice, and the post-mortem showed atrophic changes in the liver. The cases continued to increase, and in August and September, 1916, 57 cases with 16 deaths were reported.

Following the introduction of protective measures, there was a steady decline in the number of cases reported, until toxic jaundice almost disappeared among munition workers. The earlier method of filling shells involved much handling and spilling of amatol, and at one factory the heaviest incidence was among the cleaners and truckers. The powder was poured into the shell through a funnel, and then hammered down by striking the upper end of a wooden rod with a wooden mallet. At each stroke a fine gust of powder was driven up, and in consequence a thick layer of dust fell on the floors and about the workers. Later, mechanical methods were introduced for filling the shells, and other steps were taken to reduce to a minimum the actual handling of the substance. In addition, a system of exhaust ventilation was established to change the air rapidly and carry away dust. Other measures included the provision of canteens, shortening of hours, attention to the general health of the workers, and a system of alternating the employment so as to lessen the period a worker was subjected to continued exposure.

The symptoms displayed could be classified under five main headings — (1) Dermatitis, (2) an early, probably reflex, vomiting, (3) affections of the blood or blood-forming organs, (4) toxic gastritis, (5) toxic jaundice.

The parts most affected by dermatitis were the wrists, ankles, and neck. Secondary septic infection sometimes occurred. Some workers showed minute ulcers on the fingers, or definite "powder holes" in the webs between the fingers or fingers and thumbs.

Reflex vomiting was seen among new workers in the early morning, and was probably psychic in origin. This symptom yielded readily to treatment, and was not observed after the first two years.

The blood changes manifested themselves as aplastic anæmia, 15 cases of which are known to have occurred, all being fatal. This disease began with gradually increasing debility and breathlessness, or was sometimes sudden and hæmorrhagic. There was no parallelism in the incidence of toxic jaundice and aplastic anæmia.

Toxic gastritis was a widespread symptom seen in all filling factories. Pain,

<sup>1</sup> *T.N.T. Poisoning and the Fate of T N T in the Animal Body*, 1921.

nausea, aversion to food, and loss of weight, and constipation were concomitant symptoms, the patients showed marked apathy and muscular weakness, and their faces were pale, drawn, and wizened

Toxic jaundice appeared most often in the third month of employment. Sometimes there were premonitory symptoms of dizziness, fatigue, and headache, but in other cases there was no warning. Vomiting was often severe. Tenderness over the liver was occasionally elicited. Emaciation and delirium preceding death were observed. Young adults were frequently attacked, and these cases showed a high mortality.

In some instances of both toxic jaundice and aplastic anæmia there was a considerable interval between the absorption of the poison and the development of the symptoms. A boy who left a factory at the end of June and thereafter worked on a farm became jaundiced on September 1st, and died on September 9th.

**Post-mortem Appearances.**—The most marked feature in the post-mortem appearances in toxic jaundice was the great reduction in the size of the liver. In a series of seven cases described by Turnbull, the weights of the livers varied from 17 ounces to 31 ounces. Microscopically there was degeneration and necrosis of parenchyma associated with infiltration and fibrosis. The myocardium showed extensive fatty degeneration. The kidneys were enlarged, rounded, and flabby, and fatty degeneration was present. Numerous petechial hæmorrhages were present in several cases.

## ANILINE.

**Aniline** ( $C_6H_5NH_2$ ), phenylamine, or aniline oil, which forms the basis of various aniline colours, is produced commercially by reducing nitrobenzene. When pure and freshly prepared, it is a colourless oily liquid, having a peculiar odour, after a time, especially with access of air, it turns brown, crude commercial aniline contains toluidine in admixture. It is very slightly soluble in water, but is freely so in alcohol and ether. Aniline can be absorbed by the unbroken skin, as well as by the lungs and mucous membranes.

**Symptoms.**—When a poisonous dose of aniline is swallowed the symptoms appear in from five to ten minutes up to an hour or more. Nausea and vomiting usually occur, and the patient experiences a sensation of giddiness and drowsiness which deepens into coma, soon after the poison is taken the lips, face, the ends of the fingers and toes, the conjunctivæ, and the lobes of the ears become cyanotic. The respirations are laboured and are frequently slow, sometimes they are accelerated. The pulse is variable, it has been found small, frequent, and irregular, but in a fatal case recorded by Smith,<sup>1</sup> it was full and slow, sixty to the minute. The skin is cold and clammy to the touch, the breath may have the odour of aniline, the pupils are sometimes dilated and reactionless, at others they are contracted and slowly respond to alternations of light and darkness, the reflexes are sometimes present and at others absent. The blood has been found chocolate-coloured, and when examined with the spectroscope, yields a spectrum like that of methæmoglobin. Muller<sup>2</sup> records the case of a woman who swallowed about 25 c c (six drachms) of aniline, and became comatose and deeply cyanosed, a little of the blood taken from the finger gave the spectrum of methæmoglobin, and on the addition of ammonium sulphide, that of reduced hæmoglobin, thus differing from the spectrum obtained

<sup>1</sup> *The Lancet*, 1894

<sup>2</sup> *Deutsche med. Wochenschr.*, 1887.



with dinitrobenzene, which does not materially change on the addition of a reducing agent. In fatal cases the cyanosis and subnormal temperature persist, and not unfrequently convulsions occur shortly before death. In mild cases the only pronounced symptom may be the blue discoloration of the lips and face, without dyspnoea, this leads to a consideration of the cause of the cyanosis.

It appears very probable that the blue colour so universally present in cases of poisoning with aniline and other benzene derivatives is not wholly due to asphyxia caused by functionless hæmoglobin, but that some of the poison undergoes such chemical changes in the system as to yield coloured products which are largely accountable for the "cyanosis". The reasons for this supposition are — That the colour differs from that of ordinary cyanosis, that it has been observed without any indication of dyspnoea, and that in those cases in which the respiratory function is profoundly implicated and recovery takes place, the blue colour persists for a time after the breathing has returned to the normal condition. Many observers have stated that the colour was unlike anything they had ever seen in simple asphyxia, and also that it was much more pronounced. Reynolds,<sup>1</sup> in reporting a case of nitrobenzene poisoning, remarks that the intense blueness of the whole body was more marked than in any kind of cyanosis he had previously seen. Dehio<sup>2</sup> describes the colour of the skin in the case of a woman who drank ten grammes of aniline as not being at all like that of ordinary cyanosis — it was more of a lead tint, it did not produce the impression that it was due to overfilling of the veins (an explanation that has been suggested), for it remained when the blood was pressed out with the finger, the appearance resembled that due to a pigment which had transuded from the blood within the vessels into the skin. An interesting incident, illustrating the production of blue discoloration without dyspnoea, is related by Rayner<sup>3</sup> — A kind of epidemic in newly born children broke out in a work-house, in typical cases the lips, gums, and palate were deep blue, and the entire surface of the body was dusky, although the children were quite lively, the breathing was natural and the temperature normal. It was discovered that napkins, name-marked with a large four-and-a-half inch oval stamp charged with aniline chloride, were being used without being previously washed. The buttocks and the vulvæ of the children were stained with the pigment, which had been absorbed through the skin, when the napkins ceased to be used, the children gradually regained their natural colour. A somewhat similar instance is recorded by Landouzy and G. Brouardel,<sup>4</sup> in which a number of young children became "cyanosed" in appearance, from absorption of aniline used to colour the shoes they were wearing. Persistence of discoloration, after disappearance of the conditions causative of cyanosis, is illustrated by three cases of antifebrin poisoning (*v* p 475), in which methæmoglobin, though found in the blood during the stage of cyanosis, had completely disappeared some time before the skin resumed its natural colour. In another case of antifebrin poisoning, the patient suffered no distress, although the skin was bluish-grey and continued so for over a fortnight, in this case no methæmoglobin was found in the blood. It has been denied that aniline undergoes any changes in the body which could lead to the formation of pigments, but there are strong reasons for assuming such changes to be possible. Dragendorff<sup>5</sup> investigated the case of a woman who became comatose after swallowing about three drachms of aniline, most of which was rejected by vomiting, eighteen hours after,

<sup>1</sup> *Med. Chron.*, 1889

<sup>2</sup> *Berliner klin. Wochenschr.*, 1888

<sup>3</sup> *Fortschritte d. Med.*, 1887

<sup>4</sup> *Brit. Med. Journ.*, 1886

<sup>5</sup> *Gaz. des Hôpitaux*, 1900

the tips of the fingers, the feet, lips, and gums were cyanosed. The rejected contents of the stomach gave the reactions of aniline and of paratoluidine, the urine contained only a minute quantity of aniline, but much more paratoluidine, Dragendorff believes that some of the missing aniline was converted in the body into pigment. Exceptionally, aniline may cause destruction of the red corpuscles, and, in consequence, icterus, followed by hæmoglobinuria, has been observed.

**Fatal Dose.**—Six drachms have proved fatal, probably much less might be so.

**Treatment.**—The same as in poisoning with nitrobenzene.

**Post-mortem Appearances** are not characteristic. In one case the veins were found distended with dark-coloured blood, the mucous membrane of the bronchi and of the stomach was in parts swollen and reddened.

**Chemical Analysis.**—Aniline may be separated from organic matter by making alkaline and then distilling the mixtures, if much aniline is present, it will be visible as oily looking drops in the distillate. It may also be separated from organic matter by making alkaline and shaking out with ether.

**Tests.**—An aqueous solution of aniline, treated with bleaching powder cautiously added, yields a purple colour, tending to black. A drop of aniline on a colour-slab, treated with a drop of strong sulphuric acid, yields a dirty-white solid, if this is mixed with a couple of drops of water, and then with a drop of a solution of potassium dichromate, a bronze-green colour is produced, which rapidly changes to blue and then to black. If a minute quantity of aniline is dissolved in an aqueous solution of phenol, and a solution of bleaching-powder is dropped into the mixture, a yellowish streak, which shortly changes to blue, follows the course of each drop as it passes through the liquid. The previously described test with chloroform, resulting in the formation of phenylisocyanide, may be utilised in the detection of aniline.

**Phenylhydroxylamine** ( $C_6H_5NHOH$ ), obtained by reducing nitrobenzene, is a powerful reducing agent possessing strong basic properties, it is soluble in ten parts of boiling and in fifty parts of cold water. It is an unstable substance, changing in dilute alkaline solution into nitrobenzene, and in acid solution into aniline. It is one of the most virulent of blood poisons: it speedily disintegrates the red corpuscles and converts the liberated hæmoglobin into methæmoglobin. Lewin<sup>1</sup> injected some phenylhydroxylamine under the skin of a rabbit, and in three to four minutes the red colour of the blood in the animal's ears changed to brown. When in solution it is rapidly absorbed through the skin. Hirsch and Edel<sup>2</sup> record the case of a student who accidentally broke a flask containing an alcoholic solution of phenylhydroxylamine, which saturated part of his clothing and came in contact with the skin of the abdomen and thighs. In about fifteen minutes he was comatose and pulseless, with stertorous breathing and absence of corneal and pupillary reflexes. The lips, the mucous membrane of the mouth, and the skin of the limbs were of an intense blue, and numerous reddish brown spots, which did not disappear on pressure, were observed on the hands, thighs, and abdomen. The heart's action was extremely feeble. The urine contained albumin and casts. The blood was brown in colour, and contained a large amount of methæmoglobin. The patient recovered, but his normal colour did not return until the third day. In the urine from this case Lewin found neither azoxybenzene, nitrobenzene, aniline, nor amidophenol—the decomposition products of phenylhydroxylamine—and, therefore, believes that the poison acts directly on the blood without being decomposed.

**Paraphenylenediamine** [ $C_6H_4(NH_2)_2$ ] enters into the composition of some hair-dyes, the use of which has produced toxic symptoms by cutaneous absorption. In animals the symptoms comprise injection of the conjunctivæ, chemosis, œdema of the eyelids, proptosis,

<sup>1</sup> *Archiv. f. exp. Pathol.*, 1895.

<sup>2</sup> *Deutsche med. Wochenschr.*, 1895.

and, occasionally, convulsions. Puppe<sup>1</sup> found that it tends to form thrombi in the blood-vessels, and to produce parenchymatous inflammation of the heart, kidneys, and liver, thrombi formed by the injection of paraphenylenediamine into the veins yield the spectrum of hæmatin, whilst the blood itself retains that of oxyhæmoglobin. The lethal dose is estimated at about 0.1 grm per kilo weight of the animal.

**Pyridine** ( $C_5H_5N$ ) is one of a series of basic products contained in coal-tar which are metameric with aniline. It has a repulsive odour, and, amongst other uses, serves to render alcohol (with which it is mixed to form methylated spirit) undrinkable. A fatal case of accidental poisoning in a man aged twenty nine, by "half a cupful" of pyridine, is recorded by Helme.<sup>2</sup> The symptoms were vomiting, pallor, cyanosis of lips, elevated temperature ( $104^{\circ} F$ ), quickened pulse (128), which was weak and intermittent, rapid respirations (40), with mucous râles, tightness in the chest and pain in the stomach, the breath and expectorated matter had the odour of pyridine. Congestion of the lungs with acute delirium supervened, and death took place forty three hours after the poison was swallowed. At the autopsy the larynx, trachea, and bronchi were found to be lined with a friable yellow membrane, and the lungs were congested and oedematous, the œsophagus and the cardiac end of the stomach were greatly congested. No odour of pyridine was observed when the autopsy was being made.

**Methylacetanilide**, or **Exalgin**, has several times given rise to dangerous symptoms of the aniline type. Bokenham and Jones<sup>3</sup> state that a woman, aged twenty-four, after taking exalgin in six grain doses three times a day for a week, became blue in the lips and cheeks, with a feeling of weight in the epigastrium, subsequently she became delirious, and was more deeply cyanosed—the inhalation of amyl nitrite, by causing dilatation of the vessels, temporarily increasing the cyanosis, stimulants and strychnine were administered, and recovery took place. Gilray<sup>4</sup> records a case in which seventeen and a half grains of exalgin, swallowed in mistake for sulphonal, produced unconsciousness, recurrent convulsions, profuse perspiration, foaming at the mouth, weak, rapid, and intermittent pulse, and dilated pupils, with ultimate recovery. Bell<sup>5</sup> relates a recovery after 150 grains of exalgin which produced absolute unconsciousness, lividity, full pulse, elevation of temperature, and the presence of albumin in the urine. Weber<sup>6</sup> saw a case in which recovery followed 248 grains, coma, cyanosis and epileptiform convulsions occurred, with suppression of urine for twenty-four hours, the first urine passed contained blood. In a case recorded by Crookshank,<sup>7</sup> complete unconsciousness for three hours followed a dose of only five grains of exalgin. Beorhna Nigris<sup>8</sup> found methæmoglobin in the blood after the administration of exalgin, he states that it diminishes the number of red corpuscles, the percentage of hæmoglobin, and the excretion of  $CO_2$ .

**Acetanilide**, or **Antifebrin**, has frequently produced toxic symptoms accompanied by cyanosis. Marenchaux<sup>9</sup> relates the case of an infant, five months old, in whom a little over three grains of antifebrin, accidentally given, produced deep cyanosis, coldness of the surface, and insensibility, with excessively quick breathing—72 respirations to the minute—the pulse being 160, recovery took place. Five grains have produced cyanosis and collapse in an adult. Muller distinguished the bands of methæmoglobin in the blood of three patients who were cyanosed after taking large doses of antifebrin, he noticed that the blood became normal before the cyanosis disappeared, and attributes its continuance to overfilling of the veins, neither antifebrin nor aniline was found in the urine, but there was an increased amount of combined sulphuric acid. "Headache" or "Daisy" powders, which contain from four to ten grains of antifebrin, are sold to the public. Dumsey<sup>10</sup> records the case of a woman who took six of these powders daily for a month, she suffered no distress, but the skin became bluish grey, the discoloration lasting more than a fortnight, the conjunctivæ also had a bluish hue. No methæmoglobin was detected in the blood, which, however, was distinctly darker than natural and was of a purplish tint, nor was aniline found in the urine. Severe and even fatal poisoning by the external application of acetanilide as an antiseptic to raw surfaces has occurred. Snow<sup>11</sup> relates the case of an

<sup>1</sup> *Vierteljahrsschr. f. ger. Med.*, 1896

<sup>2</sup> *Brit. Med. Journ.*, 1893

<sup>3</sup> *Ibid.*, 1890

<sup>4</sup> *Brit. Med. Journ.*, 1892

<sup>5</sup> *The Lancet*, 1899

<sup>6</sup> *La Semaine Méd.*, 1894

<sup>7</sup> *The Lancet*, 1895

<sup>8</sup> *Annali di Chim. e Farm.*, 1892

<sup>9</sup> *Deutsche med. Wochenschr.*, 1889

<sup>10</sup> *Brit. Med. Journ.*, 1896

<sup>11</sup> *Arch. of Pediatrics*, 1897.

infant that became cyanosed and collapsed after its unhealed navel had been dusted with acetanilide, and quotes a number of other similar cases.

Gordimer<sup>1</sup> records two cases of chronic poisoning from acetanilide "habit." One was a woman of thirty who at first denied any drug habit, but was subsequently found to have been taking for four years a daily dose of from fifty to seventy five grains of acetanilide. She complained of great weakness, palpitation, attacks of faintness and shortness of breath. The lips, ears, fingers, and toes were cyanosed and the skin of the whole body was bluish-black. The heart was dilated, and there was a murmur of mitral incompetence. The spleen and liver were enlarged and tender. An erythrocyte count showed 9,200,000 per cubic millimetre with some poikilocytosis. The urine was brownish-red when passed, changing after to inkly black, the colour being due to urobilin. It reduced Fehling's solution, and yielded dextrosazone crystals with the phenylhydrazine test. The ethereal sulphates were greatly increased. Rapid and continuous improvement began when the taking of the drug was stopped. The second case, a woman of fifty two, showed similar symptoms.

Arcuili<sup>2</sup> reports the case of a girl, aged two, who took 30 "anti-influenza" tablets, each containing one grain of acetanilide. Her brother, aged three, got hold of a box of the tablets, and, having sucked off the sugar coating, gave what remained to the girl, who swallowed them. Severe prostration followed, but recovery occurred.

**Chemical Analysis**—Antifebrin may be extracted from acid aqueous solution by ether or chloroform.

**Tests**—Sulpho vanadic acid produces a brownish red, which changes to dirty green. If a drop of a solution of potassium bichromate is mixed with a drop of strong sulphuric acid on a colour slab, and a fragment of antifebrin added, a red colour which changes to brown and then to dirty green is produced. Boiled with an aqueous solution of potash, antifebrin is decomposed into aniline and potassium acetate, which may be recognised respectively by appropriate tests. Antifebrin may be distinguished from antipyrin by the absence of reaction on the addition of free chloride.

**Phenyldimethylpyrazolone, or Antipyrin**, has occasionally produced toxic symptoms. Rapin<sup>3</sup> reports the case of a woman, aged twenty eight, who took four doses of fifteen grains each on five consecutive days without ill effect, on the sixth day she took a like dose, which produced collapse, the cheeks and lips were almost black from "cyanosis", and a rash came out on the body. She was well on the following day. Blakeney<sup>4</sup> records a case in which ten grains of antipyrin immediately produced a burning sensation in the mouth and throat with acute discomfort in the abdomen, followed in five minutes by severe vomiting. Rapid swelling of the lips and face, urticaria, with collapse and imperceptible pulse, occurred. Recovery took place, but the swelling of the face did not subside for three days. Wallace<sup>5</sup> saw a healthy girl of twenty who had taken ten grains of antipyrin. Within fifteen minutes she complained of feelings of faintness and suffocation, and her face became "blotchy" and swollen. When seen, she was perspiring freely, the skin of the neck and face was oedematous, and there was profuse urticarial eruption on the face, neck, and upper part of the chest. The pulse was 120, very feeble and irregular, and temperature 95.5° F. Collapse followed with cyanosis and unconsciousness. The symptoms disappeared in two days.

**Chemical Analysis**—Antipyrin may be extracted by chloroform from both acid and alkaline solution, but it is preferable to render alkaline before shaking out.

**Tests**—With ferric chloride a dark red colour is produced, which is destroyed by mineral acids in excess. When antipyrin is heated with a solution of bleaching powder a brick red precipitate is formed. If a little potassium nitrite is dissolved in water, and excess of strong sulphuric acid is added, the nitrous acid set free produces a green colour with antipyrin, this test is common to all pyrazolones. The urine from patients taking antipyrin yields the ferric chloride reaction on simple addition of the reagent. Antipyrin is precipitated by most of the alkaloidal group reagents.

**Para-acet-phenetidin, or Phenacetin**, on several occasions has produced untoward symptoms, and in two instances death. Fraenkel<sup>6</sup> relates that a strong man after taking 108 grains of phenacetin became blue black in the face and the mucous membranes, the respirations being reduced to from two to three per minute, recovery took place. He also mentions the case of a girl, aged seventeen, who died in a few hours after taking fifteen grains of phenacetin. Betts<sup>7</sup> saw a man who took eight grains of phenacetin every three

<sup>1</sup> *Boston Med and Surg Journ*, 1911

<sup>2</sup> *Lancet*, May, 1921

<sup>3</sup> *Revue Méd de la Suisse rom*, 1888

<sup>4</sup> *Brit Med Journ*, 1899

<sup>5</sup> *The Lancet*, 1910

<sup>6</sup> *Ver eins-Beilage der Deutsch med Wochenschr*, 1895

<sup>7</sup> *Brit Med Journ*, 1896

hours, after the third dose he felt very ill, the face was dark, almost mahogany coloured, and there was shivering, inspiratory dyspnoea, and profuse sweating on the forehead, the other parts of the body being dry, on the exposed parts the skin was elevated in wheals, the temperature was below 95° F. The next day the patient was quite well. Krong<sup>1</sup> relates the case of a boy, aged seventeen, who took four fifteen-grain doses of phenacetin in three weeks and finally a fifth dose, soon after he was seized with vomiting, diarrhoea, and headache, he was "cyanosed," and the urine became chocolate coloured and subsequently contained blood, death occurred in three days after the last dose was taken. Cyanosis, methæmoglobin, vertigo, and ataxia have occurred after medicinal doses of the drug.

**Tests**—Sulpho vanadic acid produces an olive green coloration which becomes black on warming. When a solution of phenacetin is heated with some sodium persulphate a yellow coloration is produced which becomes orange coloured on prolonged boiling.

**Naphthalene** ( $C_{10}H_8$ ) is a coal tar derivative which occurs as colourless, crystalline plates, having a faint odour, it is insoluble in cold water, slightly soluble in hot, and freely soluble in alcohol and ether. It is used in medicine as an antiseptic, and when taken internally is supposed to act on the mucous membrane of the bowel without being absorbed, but even when quite pure it may produce toxic effects. Rossbach<sup>2</sup> relates a case in which toxic symptoms occurred after six grammes, the lips and cheeks were slightly cyanosed, and twitchings of the muscles occurred over the whole body, the urine was dark brown, and became inky black on standing. Zangerle<sup>3</sup> saw a boy aged twelve who had swallowed four grammes of naphthalene. He appeared as though he was under the influence of alcohol, he had a rickling gait, and could not answer questions. He was drowsy for the next four days. There was no nerve symptom, nor was the urine discoloured. Recovery took place. Severe symptoms of poisoning have been produced by seven grams. Cases have occurred in which naphthalene has produced hæmoglobinuria and strangury. Symptoms of poisoning have also followed its external use. An atmosphere charged with the vapour given off from naphthalene is capable of causing chronic poisoning. Lutz<sup>4</sup> records three cases in which symptoms of poisoning were caused by the use of naphthalene in rooms in order to preserve books, manuscripts, and linen from the ravages of insects. The symptoms comprised headache, mental depression, digestive disturbances, with irritation of the bladder, and olive green coloration of the urine. Sometimes the symptoms are very persistent, notwithstanding the removal of the patient from the contaminated atmosphere.

**Resorcin** ( $C_6H_6O_2$ ) in toxic doses produces symptoms analogous to those due to phenol. Murrell<sup>5</sup> records the case of a girl, aged nineteen, in which two drachms produced almost immediate giddiness, and a sensation of pins and needles all over the body, she became insensible, and perspired profusely, the temperature was low, the pulse imperceptible, the face pallid, the lips were blanched, the pupils normal, the conjunctivæ insensitive to the touch, the chest walls almost motionless, and there was a state of general muscular relaxation, recovery took place. In another case epileptiform convulsions occurred.

**Tests**—Ferric chloride produces a violet colour, and sulpho vanadic acid blue and then violet. If a crystal of sodium nitrite is mixed with a drop or two of concentrated sulphuric acid and a little resorcin is added, a violet colour, which changes to blue and then brown, is produced.

**Pyrogallol** ( $C_6H_6O_3$ ), or pyrogallie acid, when absorbed into the system in large amount, destroys the red corpuscles, causes dyspnoea, reduction of temperature, lessened sensibility, paralysis, and the presence of free hæmoglobin and methæmoglobin in the urine, methæmoglobin has been found in the blood. The dyspnoea, which may be excessive, is probably due to the formation of thrombi, which are the ultimate cause of death.

Four cases of fatal poisoning due to external application have occurred since the introduction of pyrogallol in 1878 in the treatment of psoriasis. When extensively applied to the surface in the form of an ointment absorption takes place, the following is a *résumé* of the results which occurred in all four cases.—The toxic symptoms came on very suddenly in one case after the first rubbing, in a second case on the third day, in a third case on the sixth day, and in a fourth on the fifteenth day, they comprised rigors, nausea, prostration, quick pulse, primary elevation followed by rapid fall of temperature, acute anemia, jaundice, vomiting, diarrhoea, albuminuria, hæmoglobinuria, hæmaturia, broncho pneumonia, and great dyspnoea. On section the kidneys were black and intensely congested, and the blood was black and fluid.

<sup>1</sup> *Berliner klin. Wochenschr.*, 1895.

<sup>2</sup> *Therap. Monatsschr.*, 1899.

<sup>3</sup> *Berliner klin. Wochenschr.*, 1884.

<sup>4</sup> *Verhandl. d. Soc. Scientif., S. Paulo*, 1906.

<sup>5</sup> *Med. Times and Gazette*, 1881.

Reilly<sup>1</sup> records the case of a woman, aged thirty-two, who swallowed half an ounce of pyrogallol. When discovered the face was grey, the lips, cheeks, and ears being dark blue, vomiting, diarrhoea, and hæmaturia occurred, but there was no abdominal pain. The heart's action was depressed from the first, and the patient became comatose and died sixty-eight hours after the symptoms were first noticed. Post mortem, all the viscera were congested and the kidneys were dark purple in colour. Dalche<sup>2</sup> states that a man, aged twenty-three, who swallowed rather less than half an ounce of pyrogallol in solution, immediately felt a burning sensation in the œsophagus and stomach, followed by persistent vomiting. On the following day methæmoglobinuria and hæmoglobinuria occurred, the patient had cramps in the arms, became comatose and died. On section, no changes were found in the digestive tract, brain, or lungs, the kidneys were swollen and blackish in colour, granular fragments were found in the tubules and coagula in the veins, the spleen was enlarged and contained granules like those in the kidney. Benerj<sup>3</sup> records a case in which a man and his wife each swallowed more than a drachm, vomiting was procured in half an hour, the man alone experiencing further symptoms, which were limited to a sensation of drowsiness coming on at intervals, nausea, slight paroxysmal numbness about the limbs and face, palpitation, dryness of the throat, and a black tongue, the following day he was quite well. A quarter of an ounce of pyrogallol in solution has proved fatal to an adult. Petrone<sup>4</sup> records the poisoning with pyrogallol of five persons in one household, two of them died on the seventh day, the others recovering slowly. He also gives the results of a number of experiments that he made on animals for the purpose of ascertaining the toxic action of pyrogallol. A case of mixed poisoning is related by Maillart and Andeoud<sup>5</sup> in which a man swallowed about four grains of pilocarpine, and, immediately after, two drachms of pyrogallol, in three or four minutes most of the poison was rejected by vomiting. The symptoms that ensued were due to the pilocarpine, profuse sweating occurred, with increased secretion from all the mucous surfaces and the glands, pain and tenderness in the abdomen, reduction of temperature to 96° F, and temporary abolition of sight, which was quickly restored by atropine, recovery took place. Both pilocarpine and pyrogallol were detected in the urine.

**Treatment** consists in evacuation of the stomach if necessary, the administration of stimulants, the inhalation of oxygen, and external warmth.

The **post-mortem appearances** are not characteristic.

**Chemical Analysis**—Pyrogallol may be dissolved out of dried organic matter by digestion with alcohol. After filtration the alcohol is evaporated, the residue is extracted with water and shaken out with ether, which on evaporation leaves the pyrogallol behind.

**Tests**—With lime water a purple red is produced, with basic lead acetate a reddish colour, and with ferrous sulphate a bluish black. A solution of sodium molybdate added to a solution of pyrogallol produces a brownish red colour.

**Salicylic Acid** ( $C_7H_6O_3$ ) used in medicine for the most part in combination with sodium, has occasionally produced toxic effects. The symptoms vary, they comprise—hæmorrhage from the gums and the kidneys, retinal hæmorrhage, epistaxis, hæmaturia, albuminuria, vomiting, irregularity of the pulse, urticaria, hallucinations, and insensibility. Parenchymatous nephritis is a constant result of salicylate poisoning in both animals and human beings. Vinci<sup>6</sup> records the case of a man who died in thirty-one hours after taking rather more than one ounce of sodium salicylate, the autopsy revealed parenchymatous nephritis. Charters and Maclellan<sup>7</sup> state that the toxic effects of sodium salicylate are due to impurities which exist in the artificially prepared salt, and that the natural salt is not toxic. In one of two cases recorded by Auld,<sup>8</sup> one hundred grains daily for six days caused great dyspnoea and stridulous breathing, extreme slowness of the pulse, general paralysis, and some delirium, in another case the delirium was more marked. Recovery has occurred after very large doses. A patient was given an ounce and a half of sodium salicylate in mistake for sodium sulphate, the symptoms produced were—burning sensation in the throat and stomach, thirst, nausea, vomiting, profuse sweating, coldness of the limbs, defective vision without alteration of the pupils, slow action of the heart, noises in the ears with deafness, and a state of collapse, which with the deafness lasted several days, albumin was present in the urine.<sup>9</sup> Koelin<sup>10</sup> states that a man, aged twenty-two, after taking about seventy-five grains of salicylic acid in seven hours, developed ringing in the ears,

<sup>1</sup> *Brit Med Journ*, 1897

<sup>2</sup> *La Semaine Med*, 1896

<sup>3</sup> *The Lancet*, 1892

<sup>4</sup> *Ricerche clin experiment dell' avelen da acido pirogall*, 1895

<sup>5</sup> *Revue Méd de la Suisse rom*, 1891

<sup>6</sup> *Arch di Farmacol sper*, 1905

<sup>7</sup> *Glasgow Med Journ*, 1889

<sup>8</sup> *The Lancet*, 1890

<sup>9</sup> *Deutsche med Wochenschr*, 1881

<sup>10</sup> *Corresp Blatt f Schweiz. Aerzte*, 1896

deafness, stammering speech, loss of power of deglutition and of consciousness, with violent attacks of mania. Four days after the breathing was very slow and of the Cheyne Stokes type, the pulse was small and frequent, the temperature was subnormal, the pupils were contracted, and the face, neck, and thorax were cyanosed. The urine was green and contained salicylic acid, blood, and albumin. It was afterwards found that the drug administered was chemically impure.

**Chemical Analysis**—From organic matter salicylic acid may be separated by acidulation and then shaking out with ether.

**Tests.**—Both salicylic acid and phenol strike a violet colour with ferric chloride, the addition of acetic acid destroys the colour produced by phenol, but leaves that produced by salicylic acid unaltered. In ammoniacal solution salicylic acid undergoes no change on the addition of bromine water, under like conditions phenol turns blue. Salicylic acid is chiefly eliminated by the kidneys and may be detected in the urine by the addition of ferric chloride.

**Aspirin, or Acetyl-salicylic Acid**, is a white, almost insoluble powder largely used in medicine as an analgesic. Several cases are recorded in which comparatively small doses have produced serious toxic symptoms. Kirkman<sup>1</sup> took ten grains for headache, and shortly afterwards felt tingling in the arms and feet, and a tendency to retch. He experienced a burning sensation over the whole body, which became covered with an urticarial rash. The fingers and toes, and face were swollen, the lips double the normal size, the conjunctivæ suffused, and the eyelids puffy. The pulse became too rapid to count, and there was a brief period of unconsciousness. The symptoms gradually disappeared in twenty-four hours. Other observers have recorded similar symptoms after doses of five, ten, and fifteen grains. In all these, rapid swelling of the face and an urticarial rash were marked features. In view of the great frequency with which aspirin is prescribed and rarity of poisoning, it is probable that there was a special idiosyncrasy to the action of the drug in these cases, or that some impurity was present, but Kirkman advises that patients should always be advised to rest for an hour or two after its administration.

Lewis<sup>2</sup> reports the case of a man, aged twenty-four, who took probably nearly 200 grains of aspirin in the course of six hours. He did this in order to get fit quickly, as he was under instructions for France. On admission to hospital on October 25th, he was markedly anæmic, temperature 101.4° F, pulse 120. During the day he vomited. On October 26th anæmia was more profound, pulse 150, weak and irregular. An enema was administered with little result. Vomiting continued at intervals. On the following morning at 5 a.m. a large quantity of blood was passed by the bowel, and he rapidly became unconscious. He died a few hours later. Post mortem, the last five feet of the ileum was actually congested, and the cæcum and colon were loaded with blood clots. The line of demarcation between the healthy and congested bowel was very definite. The small intestine was uniformly inflamed, and the mucous coat had apparently disappeared, leaving the sub-mucous coat and blood-vessels exposed and eroded. Hemorrhage from this large area was the cause of death. The other organs were healthy. Lewis points out that the aceto-salicylic acid is converted in the upper portion of the small intestine into free salicylic acid, and it is probable that the salicylic acid was responsible for the removal of the lining membrane of the bowel. The mucous membrane of the cæcum and colon appeared to be unaffected.

## PHENOL OR CARBOLIC ACID.

**Phenol** ( $C_6H_5OH$ ), when pure, forms a crystalline mass without colour, which reddens if exposed to air. The change in colour is due to oxidation and not to the presence of by-products, such as cresol and the like, chemically pure phenol becomes red if frequently melted with free exposure to air. It has a penetrating odour indicative of its presence in the smallest amount. Although commonly called **carbolic acid** it has no acid reaction, but it coagulates albumin and destroys tissue, it is slightly heavier than water, in which it dissolves in the proportion of 1 part to 15, it is freely soluble in alcohol and ether. For sanitary purposes crude carbolic acid is used, which consists of from 15 to 60 per cent of phenol along with a varying admixture of other products of distillation from coal-tar, it is a dark-coloured liquid having the odour of phenol, modified by the impurities which are present.

<sup>1</sup> *Brit Med Journ*, 1911.

<sup>2</sup> *Brit Med. Journ*, Jan 1919.

When strong carbolic acid is applied to the skin it causes a white appearance, the epiderm is destroyed and easily peels off, the part subsequently becoming brown and parchment-like, absorption may take place through the unbroken skin to such a degree as to cause death, a lotion containing 5 per cent has produced severe symptoms of poisoning.

The toxic action of carbolic acid is both local and remote, locally it acts as a corrosive, remotely it exercises a complex influence on the nervous system. In animals it first stimulates and then paralyses the centres in the brain and cord, in human beings poisonous doses seem to paralyse from the first. The vasomotor and respiratory centres are early affected the pulse becomes small and of low tension, and the breathing irregular and laboured, almost at the same time the higher centres are attacked giddiness, reeling gait, and tendency to delirium are quickly followed by profound coma. In some instances the rapidity with which the cortical centres are attacked is very striking, and puts the local symptoms quite into the shade. Death is due to respiratory and cardiac paralysis.

**Symptoms.** When the strong acid is swallowed, immediate burning pain is experienced from the mouth down to the stomach, a sensation of giddiness and impending loss of consciousness is felt, soon followed by coma and collapse. The face is ghastly, the breathing is stertorous, the lips are livid, or stained and swollen from contact with the poison, the pupils are contracted, the pulse, small and scarcely perceptible, is usually rapid the temperature is low, and the surface is bedewed with moisture or it may be dry. Vomiting is not so constant as in poisoning by other corrosives it may not only be absent, but may be even difficult to procure. The urine is usually diminished or suppressed, that which is voided is often dark in colour, or becomes so on exposure to the air, probably due to an oxidation product of phenol—hydroquinone. Much of the phenol and hydroquinone that is eliminated in the urine is in combination with the sulphuric acid of the sulphates, hence when freshly voided the urine may be of normal colour, but subsequently becomes dark from the liberation and further oxidation of these products. Albumin and casts and, exceptionally, blood may be present. The remote effects of phenol have been developed by its injection into the bowel, an instance occurred in which about 144 grains, diluted with water, were administered to a boy of five, as an enema to kill worms, no pain was produced, but immediate insensibility, which terminated in death in about fourteen hours<sup>1</sup>. Death in an adult has been caused by rinsing out the mouth with carbolic acid in mistake for vinegar.

The external application of phenol has proved fatal. Warren<sup>2</sup> mentions a case in which it was applied to the back of an adult, producing coma, trembling of the muscles, and death in twenty minutes. Injection of phenol into abscess cavities has also caused death. Prolonged breathing of air impregnated with the vapour of phenol may produce symptoms of poisoning. Unthank<sup>3</sup> relates the case of a man who was exposed for three hours to the fumes of strong phenol, he was seized with giddiness, stupor, and convulsions. When seen shortly after, he was comatose, the neck and face were livid, the surface was cold, and the pulse scarcely perceptible, recovery took place.

Poisoning by carbolic acid is almost invariably either suicidal or accidental, the facility with which it may be obtained accounts for the former, and carelessness for the latter. In poorer households it may be kept in an ordinary wine bottle, which leads to its being drunk in mistake for some potable fluid,

<sup>1</sup> *The Lancet*, 1883

<sup>2</sup> *Med. Press and Circ.*, 1882.

<sup>3</sup> *Brit. Med. Journ.*, 1872



or, if contained in a medicine bottle, it is thoughtlessly given instead of physic. In 1919 the number of deaths from accidental poisoning by carbolic acid was 23, and from suicidal poisoning 38, in England and Wales.

**Fatal Dose.**—One drachm has caused death in twelve hours. Sometimes death occurs very rapidly, in less than half an hour, it has occurred in three minutes, and, on the other hand, it has been delayed for sixty hours, the usual period is from three to four hours. Recovery has followed enormous doses. Greenway<sup>1</sup> reports the case of a woman who swallowed more than an ounce of carbolic acid containing 90 per cent of phenol, there was profound collapse and total insensibility, but recovery took place. Davidson<sup>2</sup> records the recovery of a woman of forty, after she had swallowed four ounces of crude carbolic acid, the stomach-pump being used twenty minutes after the poison was taken. Hind<sup>3</sup> relates the case of a girl of seventeen who recovered after swallowing six ounces of crude carbolic acid, vomiting was provoked at once, the acid contained only about 14 per cent of phenol.

**Treatment.**—Although carbolic acid is a corrosive, the stomach should be emptied by means of a soft tube, if nothing but the ordinary stiff stomach pump tube is to hand, *great care should be exercised in its introduction* as the walls of the œsophagus are less resistant than in the normal condition. Several observers have noticed that apomorphine and other emetics failed to produce vomiting in phenol poisoning. After emptying, the stomach should be well washed out with luke-warm water, in which some magnesium sulphate, or saccharated lime, may with advantage be dissolved, in order to afford an opportunity for the phenol to combine and form an innocuous ether-sulphate. White of egg and milk may be given. Carleton<sup>4</sup> recommends acetic acid, in the form of vinegar diluted with water, as an antidote with which the stomach may be washed out, or it may be given by the mouth. Olive-oil has been recommended, but with doubtful advantage. External warmth, with stimulants such as ether administered hypodermically, or alcohol by the mouth or rectum, are of great value. If death from respiratory paralysis appears imminent, breathing should be promoted artificially.

**Post-mortem Appearances.**—Stains produced by the poison may be present at the angles of the mouth and on the chin, and its odour may be perceptible. The mucous membrane of the mouth may be softened, and whether white or ash-grey in colour, that of the œsophagus being similarly affected in parts, on account of shorter period of contact the changes in the mouth and œsophagus are not usually so well marked as those in the stomach. The peritoneal surface of the stomach may be injected, its mucous coat usually being corrugated, toughened, and of a brown colour, in parts it sometimes appears stiff and leathery as though it had been tanned, in other instances it is softened and easily detached. It has been observed to be of an ash-grey colour with small hæmorrhagic points, actual erosion is uncommon. Blood-stained mucus has been found in the stomach. The duodenum may present a similar appearance, the brown colour being sometimes limited to the summit of the *valvulæ conniventes*, in a preparation in the museum of Owens College this is well shown in the form of a series of parallel brown lines running across the bowel for fully twelve inches.

**Chemical Analysis.**—After the addition of a little sulphuric acid, separation from organic matter may readily be effected by distillation.

<sup>1</sup> *The Lancet*, 1891

<sup>2</sup> *Med Times and Gaz.*, 1875

<sup>3</sup> *The Lancet*, 1884

<sup>4</sup> *Therapeut Monatshefte*, 1906

**Tests.**—The presence of phenol in the distillate may be recognised by its giving with bromine-water a precipitate of tri-bromo-phenol ( $C_6H_2Br_3OH$ ), the precipitate is soluble in excess of phenol. If a little ammonia-water and a small quantity of bleaching-powder, or bromine-water, are added to an aqueous solution of phenol, on gently heating the mixture a blue colour is produced, acidulation after cooling changes the blue to red or yellow. The addition of a solution of ferric chloride to a solution of phenol produces a violet colour, and acid nitrate of mercury (Millon's reagent) a bright red. The former is not a delicate test, and the latter yields a similar reaction with proteids. If a weak solution of furfural is added to a solution of phenol, and strong sulphuric acid is allowed to trickle down the side of the test-tube, a red colour which changes to blue develops above the acid.

**Quantitative** estimation of the phenol present in the distillate may be made by precipitation with bromine water, the precipitate being washed, dried, and weighed—100 parts of tri bromo phenol correspond to 28.39 parts of phenol. The phenol may be liberated from its combination with bromine by treatment with sodium amalgam, and then extracted with ether, and the residue, after evaporation of the ether, tested as above described.

Combined phenol sulphonic acid in urine may be decomposed and the phenol estimated as follows.—Evaporate the urine to a syrup, extract with absolute alcohol, filter, and precipitate the alcoholic solution with oxalic acid as long as any precipitate falls, then add potassium hydride to feeble alkaline reaction and evaporate again to a syrup. Render the residue acid and distil the phenol thus set free from the potassium phenol sulphonate, the amount is estimated by conversion into tri bromo phenol.

**Creolin** is an oily, dark coloured liquid which forms a milky emulsion with water. It is derived from coal tar and contains naphthalene and phenol along with various hydrocarbons. It is used as a germicide, and is only poisonous in large doses. As recorded by v. Acheren,<sup>1</sup> a man, aged thirty, drank about nine fluid ounces of creolin which produced vomiting, unconsciousness, and clonic spasms, on the second day the spleen and liver were slightly enlarged and the conjunctivæ became yellow, the urine was dark green and contained coal tar derivatives, recovery took place. Pinner<sup>2</sup> saw a woman, sixty years of age, who drank about two and a half fluid ounces of creolin, she became comatose and pallid, with cyanotic lips, the pupils were rather small and reacted feebly to light, the odour of creolin was present in the breath. Vomiting and diarrhoea occurred, the urine was dark green in colour and gave a precipitate with bromine water, and a violet coloration with ferric chloride, recovery took place. Dinter<sup>3</sup> relates that three women simultaneously drank nearly two and a half ounces of creolin and recovered. Rosin<sup>4</sup> records a case in which a 2 per cent solution of creolin was used to wash out the uterus of a woman who had been confined, collapse and vomiting of matter having the odour of creolin took place and the patient died.

**Lysol** is a compound of cresol, phenol, and other coal tar derivatives with soap, it contains about 50 per cent of cresol. Its toxic action resembles that of phenol in a feeble degree. Its corrosive action is slight, the chief toxic effects are produced on the nervous system and on the heart. As might be anticipated, there has been recovery after large doses of lysol. Blumenthal<sup>5</sup> reports recovery after nearly an ounce had been fully absorbed. He points out that the toxic effects of lysol develop glycuronic acid in the system, which, by combining with the absorbed poison, renders it harmless. Wohlgemuth<sup>6</sup> also found large quantities of conjugated glycuronic acid in the urine. It has been stated that lysol does not darken the urine, as is the case with phenol, but Matter<sup>7</sup> states that the colour of the urine, light or dark, does not constitute a distinction between lysol and phenol poisoning. The treatment consists in washing out the stomach and in the administration of fat containing liquids such as milk. Puppe records two fatal cases.<sup>8</sup>

<sup>1</sup> *Berliner klin. Wochenschr.*, 1889.

<sup>2</sup> *Deutsche med. Wochenschr.*, 1895.

<sup>3</sup> *Therap. Monatshefte*, 1889.

<sup>4</sup> *Ibid.*, 1888.

<sup>5</sup> *Deutsch. med. Wochenschr.*, 1906.

<sup>6</sup> *Berliner klin. Wochenschr.*, 1906.

<sup>7</sup> Hofmeister's *Beitr. z. chem. Physiol.*  
u. *Pathol.*, 1907.

<sup>8</sup> *Deutsch. med. Wochenschr.*, 1906.

## PICRIC ACID.

**Picric Acid** [ $C_6H_2(NO_2)_3OH$ ], or trinitrophenol, is formed by the action of nitric acid on phenol. It consists of yellow prismatic or laminar crystals which are sparingly soluble in cold water, more soluble in hot water, and freely so in alcohol, it is somewhat soluble in ether and chloroform, but much more so in amyl alcohol. It is odourless, has an intensely bitter taste, possesses strong acid properties, and forms salts which detonate on percussion. A solution of picric acid stains objects yellow, and it has been used for this purpose in confectionery. On account of its bitter taste it has been used to replace hops in beer. Very few cases of poisoning by picric acid are recorded, and none with fatal results.

In experimenting on animals, Erb<sup>1</sup> found that picrates cause the blood to become dirty brown in colour, with the formation of distinct nuclei in the red discs and free nuclei in the serum, the white blood corpuscles show a tendency to undue numerical increase. The cause of death is heart paralysis. Elimination takes place by the kidneys, bowels, and mucous membranes.

**Symptoms.**—The following case related by Adler<sup>2</sup> illustrates the effects of a toxic dose.—A girl, aged sixteen, endeavoured to commit suicide by swallowing about 300 grains of picric acid mixed with water. Violent pain in the stomach and repeated vomiting speedily occurred, and diarrhoea soon followed, the sclera and the skin were coloured an intensely dark yellow, almost brown, the pupils were moderately dilated and reacted feebly to light, the fingers were spastically stretched and bent at the metacarpo-phalangeal articulations. The urine was ruby-red in colour, it contained neither albumin nor bile pigment, a slight sediment formed which partially consisted of brown-stained epithelium, the stools were fluid and ruby-red in colour. Both urine and fæces contained picric acid in considerable amount, traces of it were present in the urine six days after the reception of the poison. In a few days the discoloration of the skin diminished and the patient was quite well at the end of the week. Schwarz<sup>3</sup> relates the following case.—A man, aged forty-five, swallowed six and a half drachms of picric acid, immediately afterwards the stomach was washed out. The patient suffered from a burning pain in the stomach which radiated over the abdomen, yellowness of the skin, red spots on the face, headache, bradycardia, and suppression of urine. The urine which was subsequently passed was dark-red in colour and contained picric acid. Recovery took place. Karplus<sup>4</sup> found picric acid in the urine seventeen days after the poison was swallowed, in this case picramic acid and an increased amount of ether-sulphuric acid were also observed. Chéron<sup>5</sup> relates a case of poisoning from inhalation of picric acid dust, which caused discoloration of the skin, pain in the epigastrium, depression, delirium, vomiting, diarrhoea, and red-coloured urine, recovery took place. In another case symptoms of poisoning occurred from the application of about six grains of powdered picric acid to the vagina, in one hour the skin was discoloured and erythematous, and the urine was red, pain in the stomach and the kidneys with somnolence were amongst the symptoms, recovery took place, but the skin was discoloured for a week, and the erythema persisted for eleven days. A teaspoonful of picric acid has been swallowed without other ill effect than violent vomiting and purging.

<sup>1</sup> *Die Pikrinsäure*, 1865.

<sup>2</sup> *Wiener med. Wochenschr.*, 1880.

<sup>3</sup> *Wiener klin. Rundschau*, 1898.

<sup>4</sup> *Zeitschr. f. Med.*, 1893.

<sup>5</sup> *Journ. de Therap.*, 1880.

**Treatment.**—The stomach should be evacuated and well washed out, and elimination promoted by diuretics, and, if necessary, aperients, morphine will probably be required to relieve pain and cramps

**Chemical Analysis**—Organic matter should be acidulated with hydrochloric acid and digested with alcohol over a water bath, after filtration the alcoholic extract is evaporated to a syrup, taken up with boiling water, filtered, acidulated with sulphuric acid, and shaken out with ether, chloroform, or amyl alcohol Dragendorff directs attention to the fact that if chloroform or benzene is used for extraction, the solution, though containing picric acid, will be almost colourless, if ethyl or amyl alcohol is used the solution acquires a yellow tint The extract is evaporated to dryness and the residue dissolved in water and tested

**Tests.**—An aqueous solution of picric acid, gently warmed with a little potassium cyanide, changes to a deep blood-red colour Ammonio-copper sulphate yields a green precipitate with picric acid Basic lead acetate gives a yellow precipitate A piece of white silk allowed to remain a short time in a solution of picric acid is dyed yellow, the colour is not discharged by subsequent washing in water

### CREOSOTE.

**Creosote**, which chiefly consists of cresol and guaiacol, is slightly soluble in water, and is freely soluble in alcohol and ether It coagulates albumin and acts as an escharotic, when swallowed in poisonous doses it causes nausea, vomiting, abdominal pain, and diarrhoea Fatal poisoning by creosote is rare A case is reported by Marcard<sup>1</sup> of an infant who suddenly became ill and died in fourteen hours, the child's jacket was stained with yellowish spots, and there was a strong odour of creosote in the room Examined seven hours after the commencement of the symptoms, the mucous membrane of the lips, tongue, and mouth was partly red and partly grey, and showed signs of the action of a caustic, but no smell of creosote was perceptible, the child vomited and passed motions mixed with blood At the necropsy the lips and the tip of the tongue were brown and hard, various-sized erosions were found in the mucous membrane of the stomach, but no odour of creosote could be perceived, nor could any trace be obtained by chemical analysis of the viscera, the spots on the jacket, however, yielded evidence of the poison As the result of a series of experiments on animals it was found that when a minimum lethal dose of creosote is given and the animal lives some hours, the odour of the poison may entirely disappear In another case reported by Purchhauer,<sup>2</sup> a child ten days old was given from twenty-four to thirty drops of creosote, it became insensible, was convulsed, and died in sixteen hours At the necropsy, inflammation with corrosion of the digestive tract, and dark-coloured blood were found, and the odour of creosote was present An adult who was taking creosote medicinally gradually increased the dose until it reached one hundred drops, on one occasion she took a second hundred drops after the usual dose, Freudenthal,<sup>3</sup> who reports the case, saw her afterwards and found her insensible, breathing stertorously, with clenched jaws, cyanotic lips, contracted insensitve pupils, and absence of the reflexes, recovery took place On the other hand, Zawadzki<sup>4</sup> records the case of a woman, aged fifty-two, who died five days after swallowing three six-drop doses of creosote in milk After death two large erosions were found in the upper part of the oesophagus and others near the pylorus, the stomach was red and injected, and the kidneys were acutely inflamed

<sup>1</sup> *Vierteiljahrsschr f ger Med*, 1889

<sup>2</sup> *Friedreich's Blätter f ger Med*, 1883

<sup>3</sup> *New York Med Rec*, 1892.

<sup>4</sup> *Centralbl. f innere Med.*, 1894.

It is stated that creosote, unlike carbolic acid, does not cause the urine to become dark coloured, and only exceptionally does it produce nephritis. Creosote is eliminated by the kidneys, and after large doses its odour may be perceived in the urine.

The treatment of tuberculosis by creosote has led to the administration of enormous doses, apparently without injurious effects, the system is gradually trained to toleration by daily augmentation of the dose from a drop or two up to as many as one hundred or more drops. In Freudenthal's case just quoted, the patient, after recovery from the effects described, still further increased the dose until it reached two drachms and three-quarters twice a day. In fatal cases death may take place in from seven to twenty hours.

The **treatment** is the same as in poisoning by phenol.

The **post-mortem appearances** resemble those produced by phenol.

**Chemical Analysis**—Separation from organic admixtures is to be effected as is directed for phenol.

**Tests.**—Creosote may be recognised by its odour. In alcoholic solution it may be distinguished from phenol by adding a few drops of a solution of ferric chloride, a green colour is produced, which disappears on dilution with water. Phenol similarly treated gives a lilac colour which does not disappear on the addition of water.

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## CHAPTER XXXV

### ALKALOIDS AND VEGETABLE POISONS.

**Alkaloids** are basic bodies which may be considered as compound ammonias. Vegetable alkaloids are almost exclusively derivatives of pyridine, they contain carbon, hydrogen, nitrogen, and—with the exception of a few volatile alkaloids—oxygen. They are for the most part solid, crystalline, and colourless, a few, such as nicotine and conine, are liquid and volatile. Alkaloids combine with acids, the salts formed being more soluble, in water, than the free alkaloid. Unless specially qualified, the term "alkaloid" is applied to substances derived from plants or trees, basic products of an analogous constitution obtained from animal tissues are known as "animal alkaloids." Alkaloids possess certain properties in common, amongst which is that of being precipitated from solutions by substances which thus serve as alkaloidal group reagents, some of which throw down most of the alkaloids, others only a limited number, most of these reagents also form precipitates with ammonia.

**Group Reagents.**—**Phosphomolybdic acid**, which may be extemporaneously prepared by dissolving sodium phosphomolybdate with the aid of heat in water freely acidulated with nitric acid, precipitates almost all alkaloids, whether vegetable or animal, even when they are in extremely dilute solution, together with ammonium salts, and derivatives of ammonia, such as phenylamine, methylamine, and the like, it also precipitates salts of lead, silver, and mercury, unless sufficient nitric acid is present to keep the metals in solution. **Phosphotungstic acid** is another delicate group-reagent, yielding much the same

reactions as phosphomolybdic acid **Iodine** dissolved in water with the aid of potassium iodide gives a brown precipitate with most of the alkaloids **Potassio-mercuric iodide**—made by adding a solution of potassium iodide to a solution of mercuric chloride until the red precipitate first formed is just dissolved, leaving a colourless solution—produces a white precipitate with a great many of the alkaloids, if the alkaloidal solution is strong the precipitate is gelatinous, this reagent is less delicate than are those previously named, especially as regards morphine There are other alkaloidal group-reagents—as platinum chloride, picric and tannic acids, bismuth-potassic iodide, etc—but the above-mentioned are the best

**Special Reagents** are described in the sections respectively devoted to the various alkaloids, for the most part they are best applied to the solid alkaloid, obtained by evaporating to dryness a few drops of a solution which contains it

### STRYCHNINE.

**Strychnine** ( $C_{21}H_{22}N_2O_2$ ) occurs in several plants of the natural order *Loganiaceæ*, and is prepared from *Nux Vomica*, or from the *St Ignatius* bean, in both of which it is associated with brucine Strychnine forms colourless crystals, sparingly soluble in water and ether, rather more freely soluble in spirit, and much more freely so in chloroform It has an exceedingly bitter taste, which is perceptible in a solution composed of 1 part in 70,000 parts of water, it is one of the most permanent of the alkaloids, and may be detected in the putrefied remains of animals that have been poisoned by it Ottolenghi<sup>1</sup> found that on exposing strychnine to the action of saprophytic bacteria its toxic power, for a few days, is increased, after which it is progressively diminished, *B coli* diminishes it from the first, after an exposure of three months to the action of this bacterium strychnine loses half its potency Strychnine has powerful basic properties, and will neutralise the strongest acids, it may be exposed to the action of concentrated sulphuric acid for an indefinite time without undergoing decomposition The salts of strychnine met with in commerce are the sulphate, the nitrate, and the acetate

Strychnine is a constituent of certain powders used for killing rats, mice, and other vermin, those most commonly used are known as Battle's vermin killer Analyses made of these powders purchased at retail shops show that they are roughly measured out, and the percentage of strychnine is not constant, but each may be regarded as containing a fatal dose for an adult human being Butler's strychnine vermin-killer consists of flour and soot, with strychnine in much the same amount as in Battle's In some of these powders ultramarine is used as colouring agent, and as the gastric juice is sufficiently acid to destroy the colour of this pigment, if such a powder is swallowed no coloured particles may be found in the stomach after death

In poisonous doses strychnine causes general clonic spasms, which, from experiments on animals, are found to be due to increased excitability of the reflex centres in the spinal cord It appears as though strychnine had the power of lessening the resistance of the cells in the anterior cornua, both to reflex stimuli and to the reception of impulses from contiguous cells, the result being that a stimulus which, under normal conditions, would produce a response limited to the muscles supplied by the cells actually stimulated, spreads from cell to cell, and sets up universal spasm, an impulse set up in the cord travels

<sup>1</sup> *Vertheiljahrschr f. ger. Med*, 1896

wave-like in all directions, the motor cells having lost their self-inhibitory power. Houghton and Murhead<sup>1</sup> believe that in strychnine poisoning some resistance is removed to the passage of impulses between the posterior horn and the terminal fibres of the sensory nerve-roots which surround the motor cells, in other words, that the resistance in the fibres between the posterior horns and the cells of the anterior horns is diminished. They think it probable that strychnine acts neither on the terminal fibres nor on the motor cells, nor yet on the cells of the posterior root-ganglion. Verworn<sup>2</sup> states that, in large doses, strychnine paralyses the motor nerve-endings in the muscles, but that, even in the largest doses, it does not paralyse the muscle substance. The inhibitory influence of the higher centres is probably not interfered with, a striking illustration of this is occasionally seen in the human subject when under the influence of a toxic dose of strychnine.—The least external stimulus is sufficient to provoke the liberation of a torrent of motor nerve-impulses, which throws the whole of the skeletal muscles into the most violent movement, the slamming of a door, the touch of a hand, or even a current of air, will produce an attack, yet the patient, in the lull after a seizure, will sometimes ask a bystander to rub his legs in order to ease the pain, and the action is unattended by reflex spasm, this points to a certain degree of inhibitory control, which the higher centres are capable of exercising over the spinal centres, notwithstanding their instability.

**Symptoms.**—In three or four minutes, up to a quarter of an hour or longer, after a poisonous dose of strychnine has been swallowed, muscular twitchings, accompanied by a feeling of anxiety and impending suffocation, are experienced by the patient, and, immediately after, he is seized with a violent convulsion of a tetanic character. The arms and legs are stretched out, and the muscles of the trunk are hard and unyielding, then jerking movements occur, forcing the head and legs backwards and the trunk forwards, the feet being strongly flexed and the hands clenched. The clonic spasms increase in severity, and the extensor muscles are so forcibly contracted that the body is arched in the posture of *opisthotonos*—the back of the head and the heels forming the ends of a curve, of which the abdomen constitutes the most prominent part, exceptionally the body may be arched forward or sidewise. When this stage is reached the spasm for a time becomes tonic, the muscles of the chest and abdomen, including the diaphragm, are tense and rigid, and the whole frame remains arched and stiff, the pulse is very rapid and feeble, and respiration is much impeded or entirely arrested, producing marked cyanosis. The sufferer is fully conscious, and experiences the most acute physical pain, with mental anguish at the prospect of immediate death, which he feels to be imminent, when capable, he cries out for something to be done to relieve the agony which is vividly portrayed by the terror-struck face, the prominent eyeballs, dilated pupils, and cyanotic complexion. After a minute or more the muscles relax, the eyeballs cease to protrude, and the pupils regain their usual size, normal respiration recommences, the cyanosis disappears, and the pulse diminishes in rapidity. The patient lies exhausted, dreading a repetition of the spasm which sooner or later recurs, being provoked by the least external impulse, during the remission, which lasts from a few seconds up to five or ten minutes, the face, without losing its anxious appearance, looks less wildly agonised than during an attack. If the case is about to terminate fatally the spasms succeed each other in rapid sequence, and death usually occurs within two hours either from asphyxia produced by fixation of the respiratory muscles,

<sup>1</sup> *Medical News*, 1895

<sup>2</sup> *Arch. f. Anat. u. Physiol.*, 1900.

or, during an interval, from exhaustion, probably the result of the excessive expenditure of force, leading to paralysis of the nerve elements. When recovery is about to take place the attacks diminish in severity, and the intervening periods are more and more prolonged, until at last the patient, free from convulsions, is left feeble and exhausted, a condition from which he recovers in a few days, in some cases the recovery is more prolonged, but actual complications are very exceptional.

Along with the other skeletal muscles, those of the lower jaw participate in the spasm, it may be to such a degree as to clench a spoon or feeding vessel firmly between the teeth. The difference between the trismus of strychnine poisoning and that which accompanies the disease of tetanus, consists in the former being secondary to the spasms which affect the muscles of the limbs and trunk, whilst in tetanus it precedes the general spasms. In strychnine poisoning the muscles of the jaws are relaxed in the interval between the attacks, in tetanus the trismus persists during any mitigation of the general spasm. In fatal cases of strychnine poisoning death usually occurs within two or three hours, the patient being in his customary health previous to the commencement of the attack, tetanus is never so rapidly fatal, for several hours soreness and stiffness of the muscles of the face and neck precede the tetanic convulsions, and death rarely occurs within twenty-four hours, being usually delayed for several days.

In exceptional cases a much longer interval than usual elapses between the reception of the poison into the stomach and the commencement of the symptoms, two hours and more have been known to intervene. When a narcotic has been simultaneously swallowed the interval may be still further prolonged. Macready<sup>1</sup> reports a case in which a grain and a half of strychnine and two ounces of tincture of opium were taken, the symptoms of strychnine poisoning did not appear until eight hours after, the opium producing narcosis in the meantime. The opposite extreme is illustrated in a case reported by Fegan,<sup>2</sup> a man sucked an egg which, for the purpose of killing vermin, had been charged with from two to three grains of strychnine, symptoms commenced in from four to five minutes, and death took place in one hour and a half. Hunter<sup>3</sup> records a case in which the first convulsion occurred within five minutes, and Barker<sup>4</sup> one in which the symptoms began in from three minutes and a half to four minutes, after about six grains of strychnine were swallowed.

The period of survival, after the commencement of the symptoms, is also subject to variation. In Barker's case above quoted death took place within thirty minutes, in the case of Cook (*Reg v Palmer*, C C C, 1856), the interval was only twenty minutes, Christison records a case in which it did not exceed fifteen minutes. A man aged twenty-one years died in from fifteen to twenty minutes, after swallowing, on an empty stomach, between thirteen and eighteen grains of strychnine in solution. The shortest interval known was in Hunter's case, above cited, the patient, a woman of seventy, died five minutes after the symptoms first appeared. Death may be delayed beyond the two hours above stated to be the usual limit of survival, in rare instances it has not occurred until three, five and a half, and even seven hours after the poison was swallowed. Henry<sup>5</sup> records the case of a man who lived nine hours after taking from seven to ten grains of strychnine—eight and three-quarter hours after the commencement of the symptoms—treatment having been resorted to in the interval.

<sup>1</sup> *The Lancet*, 1882

<sup>2</sup> *Ibid*, 1889.

<sup>3</sup> *Med Times and Gaz*, 1867

<sup>4</sup> *Amer Journ of Med Sc*, 1864.

<sup>5</sup> *The Lancet*, 1893



An unusual amelioration of the symptoms took place four hours after the poison was swallowed, when the patient seemed so much better as to be out of danger, three hours subsequently the spasms again became more frequent, and he died from asphyxia in one of the seizures. Under exceptional conditions, as when a narcotic has been taken along with the strychnine, still longer periods of survival have been recorded. Spontaneous vomiting rarely occurs in strychnine-poisoning. Nickel<sup>1</sup> records a case in which this exceptional symptom was present.

**Fatal Dose.**—Half a grain of strychnine sulphate has caused death in twenty minutes, a little over a grain of strychnine has frequently proved fatal. Recovery has taken place after four, five, and even ten grains. Two cases are recorded in which twenty grains respectively were swallowed immediately after a meal, vomiting was at once procured, and in both instances the patients recovered. In a third case twenty-two grains of strychnine were in the stomach for two hours before vomiting occurred, yet recovery took place.

Idiosyncrasy plays an important rôle in determining the effect produced by minimum doses of strychnine, in some instances intolerance to the poison is due to exceptional irritability of the nervous structures towards the alkaloid, in others, retarded elimination may be the cause. Strychnine is eliminated in the urine, fæces, and saliva. Whilst making some investigations on this subject, the only cases in which the author failed to detect it in the urine of those to whom it was being medicinally administered, were two in which incipient symptoms of toxic action were produced—a feeling of apprehensiveness, accompanied by muscular twitchings, and involuntary jerking of the limbs. In neither of the cases was strychnine detected in the urine, although it was invariably found in other cases in which similar doses were being taken. The kidneys, for some reason, failing to eliminate any of the poison, it was eliminated by the liver, and possibly the stomach, and passed into the duodenum, where some of it would be reabsorbed, the rest being voided along with the fæces. This made the process of elimination slower, and the continued administration of the alkaloid led to its accumulation in the blood, until an incipient physiological effect was produced. Such cases are very exceptional, the rule is that speedy elimination by the kidneys takes place. Kratter<sup>2</sup> found strychnine in human urine half an hour after administration, in animals, Ipsen<sup>3</sup> found it in three to five minutes after reception. Elimination proceeds rapidly, Kratter failed to find strychnine in the urine forty-eight hours after its administration had ceased, a result with which the author's own experience agreed.

**Treatment.**—Administer chloroform by inhalation until it is possible to introduce the stomach-tube, and with its aid wash out the stomach, otherwise an emetic should be given, since spontaneous vomiting does not usually occur. After emptying the stomach the patient may be kept under the influence of chloroform, or chloral hydrate may be given. A striking instance of the value of chloral hydrate as an antagonist to strychnine is afforded by a case related by Jones<sup>4</sup>. A man swallowed two threepenny packets of Battle's vermin-killer, which produced typical symptoms of strychnine poisoning, the patient did not vomit, nor was the stomach emptied. Twenty grains of chloral hydrate dissolved in water were injected subcutaneously, followed by a second dose of twenty grains, and subsequently by ten grains more, twenty grains were also given by the mouth as soon as the patient could swallow, recovery took

<sup>1</sup> *Vierteljahrschr f. ger. Med.*, 1906.

<sup>2</sup> *Wiener med. Wochenschr.*, 1892.

<sup>3</sup> *Vierteljahrschr f. ger. Med.*, 1892.

<sup>4</sup> *The Lancet*, 1889.

place If death from asphyxia appears imminent, artificial respiration should be resorted to

**Post-mortem Appearances.**—The statements as to cadaveric rigidity are contradictory In the case of Cook, who was poisoned with strychnine by Palmer, the body was found five days after death much stiffer than is usually the case—the hands were stiff and firmly closed and the muscles strongly contracted, in other cases the rigidity has been of the ordinary type and duration, the usual interval of muscular relaxation intervening immediately after death In Hunter's case previously mentioned rigidity was not present fifteen minutes after death, nor yet three hours after, it was developed to a slight degree seven hours after death Internally there are no characteristic appearances Hyperæmia of the cerebral and spinal meninges has been recorded, and also fluidity of the blood, probably due to death occurring from asphyxia

**Chemical Analysis**—The alkaloid is to be separated from organic matter by the process described in Chapter XXVIII, and weighed if in sufficient amount Chloroform, or a mixture of chloroform and ether, are the best solvents for extracting strychnine from an aqueous solution, it is advisable to precipitate the free alkaloid in the presence of the solvent, and to shake out without delay, if allowed to become crystalline the alkaloid is much less soluble

The quantity contained in the various organs is inconstant The author analysed some of the viscera and their contents in three cases of suicidal poisoning by strychnine with the following results —In one, a grain and a half of strychnine (0.1 grm) caused death in about three hours, the stomach pump being used in the interval, the contents of the stomach yielded a trace only, the liver 0.013 grm, the urine (278 c c) 0.005 grm and one kidney a trace The second case, poisoned with a similar dose, also died within three hours, but without evacuation of the stomach, only the stomach and its contents (90 c c) were obtained, treated separately, both yielded evidence of the presence of strychnine, but not in weighable amount The third case was poisoned with a sixpenny packet and died in about two hours without the stomach being evacuated, as in the last case only the stomach and its contents (155 c c) were obtained, the stomach yielded sufficient strychnine for identification and no more, but from its contents 0.078 grm (1.2 grain) was obtained<sup>1</sup> The discrepancy between the last two cases is very striking in neither was the stomach evacuated, yet in one the contents yielded a mere trace, and from the other more than half the amount probably taken was isolated, a double packet of the vermin killer having been swallowed It is noteworthy that, although the stomach contained such a large amount of the alkaloid after death, the viscera itself yielded a mere trace The absorption rate of the stomach is much slower than that of the small intestines, this is especially the case as regards strychnine Meltzer<sup>2</sup> found that if six to ten milligrammes of strychnine are introduced into the full stomach of a rabbit clonic spasms occur within a short time, but if the pylorus be closed before the introduction of the poison—the circulation being good and the vagi intact—as much as two hundred milligrammes of strychnine may remain in the empty stomach for many hours without producing any effect Strychnine is absorbed the quickest by the pharynx, nearly as quickly by the rectum, the small intestines coming next, the œsophagus absorbs it but little better than the stomach The delay in the onset of the symptoms when an opiate has been taken along with the strychnine is probably due to quiescence of the stomach, induced by the opiate, with consequent retention of the strychnine in a feebly absorbent viscus, the symptoms only develop when some of the poison finally makes its way into the small intestine

After absorption, the blood and the liver contain the largest amount of strychnine, the firm organs—as the kidneys—yield much less The theory that the liver acts as a magazine and stores up strychnine is probably incorrect, it is more likely that the relatively large amount usually found in it is dependent upon the vascularity of the organ

As previously stated, strychnine resists the influence of surrounding putrefaction to a very considerable degree Wolff<sup>3</sup> communicates a case in which strychnine was detected in an exhumed corpse three hundred and twenty-two days after the reception of the poison Prescott<sup>4</sup> quotes a case in which it was found in the stomach, liver, and intestines in a body

<sup>1</sup> *Med Chron*, 1889

<sup>2</sup> *Journal of Experimental Medicine*, 1896

<sup>3</sup> *Einige Fälle von Strychninvergiftung*, Dissert., 1887

<sup>4</sup> *Organic Analysis*, 1887

exhumed one year and three days after death. In a case recorded by Haw<sup>1</sup> it was found nearly ten months after burial. In another case it was found six months after burial. As colliquative putrefaction in the remains of a person who has died from strychnine does not necessarily render the detection of that poison impossible, Ipsen<sup>2</sup> suggests that in cases of exhumation any transudation fluid present in the coffin, together with all impregnated portions of clothing which surround the body, in addition to the viscera, should be submitted to analyses.

**Tests.**—If a minute drop of a fluid containing strychnine is conveyed by the tip of the finger to the tongue, a characteristic bitter taste will be perceived unless the alkaloid is in very small amount, and its bitterness is masked by the presence of some substance possessing a strong, penetrating taste, this test should never be omitted when searching for alkaloids, and should precede chemical tests. A fragment of strychnine thoroughly mixed on a colour slab with a couple of drops of strong sulphuric acid undergoes no change of colour, if a few granules of manganese dioxide are then stirred in with the point of a glass rod, a blue colour which rapidly becomes purple and more gradually changes to orange-red, is produced. Lead peroxide, potassium dichromate, potassium permanganate, and other oxidising agents give the same reaction, but manganese dioxide is to be preferred on account of its slower action and freedom from any embarrassing intrinsic colour. On this ground ceric oxide has been recommended, as when pure it has little colour of its own, it is usually contaminated with didymium, however, which imparts a brownish-red hue almost as pronounced as that of lead peroxide, ceric oxide is much slower in its action than any of the other reagents mentioned. Mandelin's reagent, consisting of a drop of a solution of ammonium vanadate in strong sulphuric acid (1:200), yields the same colour reactions as manganese dioxide. If a mixture of strychnine and sulphuric acid is placed on a piece of platinum-foil connected with the anode of a voltaic couple, on touching the liquid with a platinum wire which forms the cathode the same colour reactions are produced as with manganese, by means of this, and the manganese test, the merest trace of strychnine—0.1 milligramme—may be identified. If strychnine is heated with dilute nitric acid, and a crystal of potassium chlorate is added, a scarlet colour is produced, which becomes brown on the addition of ammonia-water.

The **physiological test** may also be tried by injecting a few drops of a suspected solution into the dorsal lymph-sac of a small frog, and then placing it under a glass shade, if the solution contains but a minute quantity of strychnine, tetanic convulsions occur in a few minutes. After the convulsions have once occurred they may subsequently be provoked by rapping the shade, or the table on which the frog is placed.

## BRUCINE.

**Brucine** ( $C_{27}H_{26}N_2O_4$ ) is found associated with strychnine in *Nux Vomica* and in the *St Ignatus bean*. In water it is more soluble than strychnine, it is also soluble in alcohol and in chloroform, but not in ether. In solutions of equal strength brucine has a much more bitter taste than strychnine. Its toxic effects are similar, but its physiological action is only about one twenty fourth that of strychnine. Mays<sup>3</sup> states that with frogs the convulsions are later in coming on with brucine than with strychnine, and that even with a lethal dose they may be altogether wanting. Rothmaler<sup>4</sup> found that with rabbits, although the fatal doses of brucine and strychnine are as 32 to 1, tetanic spasms are produced by

<sup>1</sup> *The Lancet*, 1899.

<sup>2</sup> *Vierteljahrsechr f. ger. Med.*, 1894.

<sup>3</sup> *Journ. of Physiol.*, 1887.

<sup>4</sup> *Ueber die Wirkungskraft von Strychnin und Brucin*, 1893.

relatively smaller doses of brucine than of strychnine. Mice are endowed with a remarkable immunity from the action of brucine in comparison with that of strychnine, the relative fatal dose being as 140 is to 1. Brucine, not being accessible to the public, is practically unknown as a poison.

**Symptoms and Treatment, as in strychnine poisoning**

**Tests.**—Nitric acid added to a fragment of brucine produces a bright blood red colour which is destroyed by an excess of stannous chloride. If, after adding the nitric acid, the product is mixed with a little water and solution boiled, and then allowed to cool, the red colour is changed to purple on the addition of stannous chloride or of sodium thiosulphate, ammonium sulphide produces a similar but less characteristic reaction, and, if the reagent is in excess, free sulphur is precipitated. Sulphomolybdic acid or Frohde's reagent (prepared by dissolving, with the aid of a gentle heat, a centigramme of molybdic acid or of sodium molybdate in a cubic centimetre of strong sulphuric acid) gives with a fragment of brucine a pink or yellowish brown colour which changes to green or blue. Sulphovanadic acid produces a yellow, changing to orange red. A solution of ammonium selenate in strong sulphuric acid produces a pink colour, changing to yellow.

### NUX VOMICA.

The seeds of the *Strychnos nux vomica* are exceedingly hard and tough, and are too large to be swallowed whole unless with considerable effort, the powder, the extract, and the tincture of the seeds have produced toxic effects resembling those of strychnine. The symptoms are usually longer in appearing than when strychnine is taken, in one case a man swallowed about five drachms of nux vomica and was not affected for two hours, when he quickly died convulsed. Thirty grains of the powder and three grains of the extract have proved fatal. Hale<sup>1</sup> relates the case of a woman who died two hours after swallowing six drachms of tincture of nux vomica. Stevenson<sup>2</sup> records the recovery of a boy of twelve after taking about eight grains of the extract, both strychnine and brucine being detected in the urine.

### COCCULUS INDICUS.

**Cocculus Indicus**, or Levant nut, the fruit of the *Anamirta cocculus*, contains an active principle picrotoxin, along with other bases.

**Picrotoxin** ( $C_{12}H_{14}O_6$ ) is a colourless, neutral, crystalline body which does not form salts, it is not very soluble in water, but is freely soluble in alcohol, ether, and chloroform, it is odourless and has an intensely bitter taste. Picrotoxin acts as a gastro-intestinal irritant, and is a stimulant to the motor centres of the brain and cord. In small toxic doses it produces a tendency to stumble and reel—as in alcoholic intoxication—followed by stupor, in large doses it produces clonic spasms like those due to strychnine. Picrotoxin is eliminated in the urine.

Fatal poisoning by cocculus indicus is exceptional, probably not more than a dozen instances being recorded. Sozinsky<sup>3</sup> relates the case of a man, aged thirty-nine, who drank by mistake several ounces of whisky in which cocculus indicus berries had long been steeping, the mixture being intended for killing vermin. When seen an hour after, he had vomited once, he was unconscious, and had powerful general convulsions every five minutes, each convulsion, which commenced by twitching of the left corner of the mouth and a cry like that of an epileptic, lasting about two minutes with considerable opisthotonos, between the attacks there was complete muscular relaxation. The pupils

<sup>1</sup> *Brit Med Journ*, 1899.

<sup>3</sup> *Med News, Phil*, 1891.

<sup>2</sup> *Guy's Hosp Reps*, 1868.

were contracted, and the respirations were slow, but the heart was not much affected, profuse perspiration and diarrhœa were present, and death took place in three hours from exhaustion and failure of respiration. Shaw<sup>1</sup> records the case of a man who purchased, as he supposed, wild cherries, but which proved to be *cocculus indicus* berries, he put them into a bottle and filled it with brandy, and from time to time drank small doses without ill effect. One morning he drank a considerable quantity and afterwards felt dizzy and sick, he produced vomiting by tickling his throat, but a few minutes after he fell on the floor in convulsions and became unconscious, the convulsions continued for thirty minutes, when death took place, at the necropsy nothing abnormal was found except congestion, in patches, of the mucous membrane of the stomach. Swift<sup>2</sup> relates the case of a woman who died tetanically convulsed three-quarters of an hour after swallowing an alcoholic infusion of *cocculus* berries. the pupils were minutely contracted and the temperature was elevated.

Recovery has taken place even after very threatening symptoms. Dutzmann<sup>3</sup> records the case of a man, aged sixty, who crushed some of the berries and swallowed a handful, half an hour after he fell to the ground, vomited, perspired profusely, and was unconscious, the temperature was elevated, the pupils were normal in size but reactionless, the pulse was 80 and full, and the respirations were laboured and quickened. He then had convulsions, which were attended with foaming at the mouth, and cyanosis, the pulse rose to 110, recovery took place, but pain and oppression in the chest were felt for some days.

Death has taken place in consequence of the external application of the poison. Thompson<sup>4</sup> states that a child, aged six, who had prurigo of the scalp and was infested with vermin, was treated externally with an alcoholic solution prepared by infusing one pound of *cocculus indicus* berries in three gallons of alcohol, half an hour after, tetanic spasms came on, during which the pupils contracted to the smallest size, dilating in the intervals between the spasms, the spasms could be produced by touching the eyelid, they continued for six hours, when the patient died. The necropsy yielded negative results. Another child, to whom a similar application was made, also had clonic spasms, but she recovered. In all these cases the convulsive action of the poison completely eclipsed any gastro-enteric symptoms that might be present, in many ways the effects closely resembled those of strychnine, even to the reflex excitation of the spasms, and they corresponded to the action of picrotoxin on animals.

A minor degree of poisoning by *cocculus indicus*, called "hocussing," has occurred from its surreptitious administration in admixture with alcohol in order to produce a helpless condition of stupor favourable to the perpetration of robbery from the person. Formerly low-class publicans sometimes added small quantities of *cocculus indicus* to beer in order to increase the intoxicating effects of the beverage, and thus obtain for it a fictitious reputation for alcoholic potency.

**Treatment.**—Evacuate the stomach by the tube or an emetic. If clonic spasms are present chloral hydrate may be given, or chloroform administered as in strychnine-poisoning, artificial respiration may be necessary. In the minor degree of poisoning evacuation of the stomach with symptomatic treatment will probably be sufficient.

**Chemical Analysis.**—Picrotoxin may be shaken out of acid solution by ether or chloroform.

<sup>1</sup> *Med. News*, Phil, 1891.

<sup>2</sup> *New York Med Journ.*, 1897.

<sup>3</sup> *Wiener Med Presse*, 1869.

<sup>4</sup> *Med. Examiner*, Phil, 1852.

**Tests.**—Picrotoxin is not precipitated by phosphomolybdic acid, nor by a solution of iodine. It is dissolved by concentrated sulphuric acid, producing a yellow colour, which changes to black on heating. If picrotoxin is mixed with three times its weight of potassium nitrate, and the mixture is moistened with a few drops of sulphuric acid, and then excess of a strong solution of sodium hydroxide is added, a brick-red colour is produced. Picrotoxin reduces Fehling's solution.

### OPIMUM AND ITS ALKALOIDS.

**Opium.**—The inspissated juice of the *Papaver somniferum* contains a large number of alkaloids and alkaloidal substances, several of which possess powerful toxic properties. Morphine is the alkaloid to which opium owes its potency as a poison, next in importance come narcotine and codeine, which act as hypnotics, but are much less powerful than morphine. Thebaine, another alkaloid, and apomorphine, a derivative of morphine, act in a totally different way, the first is a convulsive and the second an emetic. With the exception of morphine, the alkaloids of opium rarely come under the notice of the toxicologist, but one other substance—meconic acid—on account of its constant presence in opium and its characteristic reactions, is frequently sought for.

The following are the more important official preparations of opium and morphine, with their strengths.—*Extractum Opi Liquidum*, 1.75 per cent of morphine, *Pilula Plumbi cum Opi*, 1 of opium in 8, *Pilula Saponis Composita*, 1 of opium in 5, *Pulvis Ipecacuanhæ Compositus*, 1 of opium in 10. *Tinctura Opi* or *Laudanum*, 1.05 per cent of anhydrous morphine. *Nepenthe* is a non official preparation of opium, one third less strong than *Tinct. opi*.

**Morphine** ( $C_{17}H_{19}NO_4$ ) is a colourless crystalline substance, which has a bitter taste and an alkaline reaction. It is slightly soluble in cold water, more soluble in hot water and in ethyl alcohol, and freely soluble in amyl alcohol, especially when hot, it is also very soluble in acetic ether, in ethyl ether and chloroform it is but feebly soluble. The salts of morphine are freely soluble in water and spirit.

### Acute Poisoning by Opium and Morphine.

**Symptoms.**—Excitation of the higher nerve-centres is the first result of the reception of a poisonous dose of opium or of morphine, with opium it occurs in from half an hour to an hour after the dose is swallowed, with a salt of morphine in solution the interval is less—from a few minutes to a quarter of an hour. The excitation may show itself by producing accelerated action of the heart, flushing of the face, and a feeling of increased mental activity accompanied by exhilaration of spirits, or it may simply cause physical restlessness, its duration is short, and it is succeeded by an opposite condition of depression of the nerve-centres. A sensation of lassitude, oppression in the head, giddiness, and a strong desire to sleep steal over the patient, who becomes more and more drowsy and less capable of responding to external stimuli, before this stage is reached the pupils are contracted, the stupor subsequently deepens into profound coma. In the earlier stages of insensibility the patient may be partially roused by being shaken and loudly spoken to, when the comatose condition is reached no external stimulus evokes any response. The muscles are relaxed, the surface is cold and moist, the features are shrunk and pallid, or cyanotic, the pupils are exceedingly contracted, the pulse is slow and compressible, and

the breathing is laboured, irregular, and stertorous, although the patient now presents all the appearance of a dying person, recovery may take place. If the case goes on to a fatal issue, the breathing becomes more embarrassed and may assume the Cheyne-Stokes type, mucous râles are heard, the pulse becomes more irregular and scarcely to be felt, the cyanosis deepens, and the face looks still more ghastly, the jaw drops, and but for the hard-drawn breaths, the appearance is that of a corpse. Twitching of groups of muscles is often observed, and at this final stage the pupils may be dilated—death is then imminent. The heart may continue to beat for a short time after respiration has ceased.

**To epitomise** in sequence the effects of a poisonous dose of opium—After a fugitive preliminary excitation the mental energy is progressively lowered until it fades into unconsciousness, voluntary movements are paralysed, the spinal reflexes are abolished, the respiratory centres succumb, and lastly, the heart fails. In the early stage the respirations are accelerated and the pulse is quick and small, later the respirations are slow and stertorous and the pulse is slow and full. The **usual interval** between the reception of the poison and death is from **six to twelve hours**.

Certain other symptoms may be present. If opium or the tincture has been taken its odour may be perceived in the breath. Vomiting may occur, and very exceptionally relaxation of the bowels, almost invariably the opposite condition—constipation—obtains. A tendency to closure of the pylorus exists, due to central paralysis of the motor fibres of the vagus (Baas)<sup>1</sup>. The urine and the saliva are suppressed in the later stages, sometimes the urine is simply retained from paralysis of the bladder, but the low blood-pressure and the excessive perspiration leave little for the kidneys to do, so that the quantity of urine is lessened, although from accumulation the bladder may be full. The urine is usually alkaline and is deficient in chlorides. The only secretion which is not diminished is the sweat, and this is usually increased from first to last.

Amongst symptoms of an *exceptional* character are—dilatation of the pupils in the earlier stage, apart from that which may occur immediately before death, acceleration of the pulse in the late stage, which has been known to alternate with, or to replace, the usual slow pulse, spasms or convulsions of a tetanic character, which are less unfrequent in the case of children than in adults. The action of opium on the cerebral cortex is to increase its motor irritability, this, so far as the evidence afforded by faradic stimulation goes, has been proved by Unverricht<sup>2</sup>. Direct stimulation of the cortex, which under normal conditions would determine the occurrence of simple motor impulses, liberates them so freely when it is under the influence of opium as to give rise to convulsive movements, the condition produced by opium is the opposite of that due to ether, chloroform, and chloral hydrate, by which the cortical irritability is lowered. In children the nerve-cells have not fully acquired their self-inhibitory capacity, it is therefore easily broken down during a state of increased irritability, in adults opium poisoning, accompanied by trismus and general spasms of a clonic, strychnine-like character, is an *extremely rare occurrence*.

A morphine *habitué*, aged forty-three, swallowed, within twenty-four hours, 576 grains of morphine hydrochloride, of which he swallowed 200 grains within two hours of his death. The symptoms produced were of a tetanic character, and the usual comatose condition did not develop.<sup>3</sup>

<sup>1</sup> *Deutsch Arch. f. klin. Med.*, 1904.

<sup>2</sup> *Centralbl. f. klin. Med.*, 1891, 1892

<sup>3</sup> *The Lancet*, 1906

In *very exceptional instances* morphine produces profound coma within a few minutes after it is swallowed, followed by death in forty minutes or an hour, in other exceptional cases the advent of the symptoms is delayed for two or even three hours, and death may not take place for more than twenty-four hours.

A noteworthy condition has been observed in severe cases of opium poisoning, partial recovery from the urgent symptoms has taken place to such a degree as to remove all apprehension, and then, after several hours' interval, the patient has relapsed into coma and died, as suggested by von Boeck,<sup>1</sup> it is probable that renewed absorption of the poison takes place under the influence of increased blood-pressure. In other instances patients who recover from the immediate symptoms succumb after a much longer interval, possibly of several days, in such cases the fatal issue is probably as much due to cardiac disease as to the poison. When recovery takes place it is, as a rule, complete, but in rare instances sequelæ have been recorded. albuminuria was present on the third day after the acute symptoms in a case recorded by Oliver,<sup>2</sup> and at about the same period in a case recorded by Huber.<sup>3</sup> A unique case is reported by Scheiber,<sup>4</sup> in which acute poisoning produced by the subcutaneous injection of morphine was followed by psychical disturbance, aphasia, and by the formation of bed-sores.

**Fatal Dose—Adults.**—Four grains of opium in one case, and two drachms of the tincture in another proved fatal. Recovery has taken place after three ounces of the tincture, equal to ninety-nine grains of opium (Burgess<sup>5</sup>), and as recorded by Bowstead,<sup>6</sup> even after eight ounces of laudanum were swallowed by a woman aged thirty-eight, who was not discovered for fourteen hours after she took the poison. One grain of morphine hydrochloride has caused death. recovery has taken place after thirty, thirty-six, and in one instance fifty-one grains, of which the greater part remained in the stomach thirteen hours (Morse). In a case reported by Bonjean<sup>7</sup> a young man swallowed fifty-five grains of morphine acetate in solution the stomach not being evacuated for upwards of two and a half hours. most dangerous symptoms followed, but recovery took place. Recovery has taken place after the hypodermic injection of probably about twelve grains of morphine (Pope)<sup>8</sup>. On the other hand, severe toxic symptoms have followed considerably less than the maximum pharmacopœial dose. Mandl<sup>9</sup> reports the case of a man who, ten minutes after a hypodermic injection of one-sixth of a grain of morphine hydrochlorate, suddenly became convulsed and then profoundly insensible with cyanosis, Cheyne-Stokes breathing followed. After four hours' continuous treatment the patient recovered consciousness, but he remained amnesic till the following day. Disease of the kidneys, especially cirrhotic kidney, predisposes to a fatal issue from opium or morphine.

**Infants.** It is well known that infants are susceptible to the influence of opium to an extraordinary degree, on more than one occasion a single drop of laudanum has been recorded as fatal. A dose of paregoric equal to the ninetyeth of a grain of opium, and in another case a dose of Dalby's carminative equal to half a minim of laudanum, are stated to have caused death, as regards such very minute doses as the two last named, there is reasonable ground for doubt, since the preparations concerned are made from crude opium, containing

<sup>1</sup> *Ziemmsen's Cyclop.*, Bd 17

<sup>2</sup> *Gaz des Hôpitaux*, 1871

<sup>3</sup> *Zeitschr f klin Med*, 1889

<sup>4</sup> *Ibid*, 1888

<sup>5</sup> *Dublin Journ of Med Sc*, 1892

<sup>6</sup> *The Lancet*, 1873

<sup>7</sup> *Annales d'Hygiène*, 1845

<sup>8</sup> *The Lancet*, 1894

<sup>9</sup> *Wiener med. Wochenschr.*, 1899



an unknown percentage of morphine Recovery in an infant three months old is recorded by Bramwell,<sup>1</sup> after a teaspoonful of laudanum (vomiting occurring soon after it was swallowed) Chamberlain<sup>2</sup> witnessed the recovery of an infant, six days old, which had swallowed a powder containing a grain and a half of opium, two hours afterwards it was apparently dead, breathing having ceased, artificial respiration was kept up for three hours, and in twenty-four hours the child was quite well Morgan<sup>3</sup> saw an infant, one month old, which was comatose after three drops of laudanum, and the breathing gradually ceased, artificial respiration was kept up almost continuously for three hours, and recovery took place after complete unconsciousness had lasted for forty-five hours Fotheringham<sup>4</sup> records the recovery of an infant, three months old, after one fluid drachm of the pharmacopœial solution of morphine hydrochlorate Egan<sup>5</sup> reports recovery in an infant seven months old, after one grain of morphine hydrochloride, vomiting, induced by emetics, did not commence until two hours after the poison was swallowed Electricity and artificial respiration were persistently resorted to for more than seven hours The infant was quite lively the following day

Preparations of opium have caused death from external application, but probably not without the skin being broken, morphine sprinkled on an open sore has proved fatal

**Treatment.**— If the poison has been swallowed the stomach-tube should be used, and the stomach well washed out, in default, an emetic may be administered by the mouth or apomorphine may be injected subcutaneously Persevering attempts should be made to rouse the patient by external stimulation, the faradic current, applied to various parts of the body with a wire brush, is an efficacious stimulant In the *less severe* forms of poisoning walking the patient to and fro between two assistants may be resorted to in such cases the cold douche and perambulating the patient may be sufficient but the former should never be used when the surface is cold, nor should the latter be carried to excess, so as to exhaust the strength *In severe cases it is worse than useless to drag a comatose individual about* When the coma is profound, artificial respiration may be needed, this constitutes *a most valuable aid to recovery* it may be supplemented by faradic stimulation of the phrenes, and, if there be much cyanosis, by the inhalation of oxygen Ammonia may be applied to the nostrils in the form of smelling-salts the vapour of ammonia-water should not be used, it is too irritating to the respiratory mucous membrane Hot coffee may be given by the mouth if the patient can swallow, if not it may be administered by the stomach-tube or as an enema Moor<sup>6</sup> recommends the administration of ten to fifteen grains of potassium permanganate dissolved in six or eight ounces of water, to be repeated three or four times at half-hour intervals, if opium or the uncombined alkaloid has been taken, the solution of permanganate should be acidulated with a little sulphuric acid Moor found that potassium permanganate oxidises morphine even in the presence of other organic matter This has been confirmed by Luff,<sup>7</sup> who found that when three grains of morphine acetate were mixed with six ounces of vomit, and the mixture was treated with four grains of permanganate dissolved in four ounces of water no morphine could be extracted from it As shown by Thornton's and Holder's<sup>8</sup> experiments on dogs, it is useless to inject this antidote hypodermically but,

<sup>1</sup> *Boston Med Journ*, 1887

<sup>2</sup> *The Lancet*, 1889

<sup>3</sup> *Boston Journal*, 1858

<sup>4</sup> *Brit Med Journ*, 1898

<sup>5</sup> *Med Times and Gazette*, 1870

<sup>6</sup> *Med Record*, 1894

<sup>7</sup> *Brit Med Journ*, 1896

<sup>8</sup> *The Therapeutic Gazette*, 1898

after poisoning by hypodermic injection of morphine, Luff recommends that at intervals the stomach should be washed out with a weak solution of permanganate in order to oxidise any of the poison which may be excreted into it. Potassium permanganate should not be administered in very concentrated solution, since it may act as an irritant, or even as a corrosive (See p 414.) One-twelfth of a grain of atropine sulphate, injected hypodermically, is recommended for the purpose of stimulating the respiratory centres, but its utility is doubtful, notwithstanding the number of successful cases in which it has been used (See section on Antagonisms of Poisons.) Hypodermic injections of ether are efficacious. Strychnine is strongly advocated by some. Lucatello<sup>1</sup> had a case in which a patient swallowed about forty-five grains of opium and twenty-two grains of morphine sulphate on an empty stomach notwithstanding which symptoms did not appear for an hour. Breathing having nearly ceased, artificial respiration, with faradisation of the phrenics, was resorted to, but without effect, under the influence of hypodermic injections of strychnine respiration was resumed.

**Post-mortem Appearances.**—Apart from discovery of the poison in the body, the post-mortem indications are not characteristic. When opium itself has been swallowed, its odour may be perceptible in the stomach, if the organ has been well washed out with the tube, or cleared with emetics, or if morphine was taken, this indication will be absent. Injection of the gastric mucous membrane has been described, it is by no means constant, and, when present, is probably due more to the treatment than to the poison. Hyperæmia of the brain and its membranes is not unfrequent, and in addition there may be œdema into the subarachnoid space and the ventricles. The amount of blood in the lungs varies; they may or may not present the appearance associated with death from asphyxia. The blood has been found dark and fluid, it has also been found coagulated.

**Chemical Analysis.**—The difficulty which attends the isolation of morphine from the organs and tissues of those poisoned by it has led to the supposition that it undergoes decomposition in the living organism. Some investigators state that it is eliminated as such in both urine and feces, others have failed to find it in the urine, but have detected it in the feces, others again have found oxidation products—as oxymorphine—in the urine, and hold that morphine is entirely changed in passing through the body.

When the delicacy and the distinctiveness of some of the reactions of morphine are taken into consideration, the undoubted difficulties that surround its detection in the organs of those who have succumbed to its influence require explanation. Neglecting for the moment the question of decomposition of morphine within the organism, there are one or two errors of procedure which may prevent its recognition. One is excess of acid in the fluid used to extract the alkaloid from organic admixture, another is the use of too high a temperature for the purpose of evaporating the solution obtained, when these two adverse conditions are combined, the probability is that small amounts of morphine which originally may be present will be decomposed and rendered incapable of recognition by the usual tests. Further, delay in shaking out after alkalisatation, and the use of an inappropriate solvent, such as ether, impedes or prevents extraction, if the aqueous solution is over alkalisated with sodium or potassium hydroxide, morphine is redissolved, and minute amounts cannot be shaken out by any solvent. The only solvents that can be depended upon to take up morphine from neutral or slightly alkaline aqueous solutions are—amyl alcohol (preferably hot), meta cresol, and acetic ether. As a solvent amyl alcohol is the best, but it is disagreeable to work with, its evaporation demands a relatively high temperature, and, as met with in commerce, it is liable to contain resinous substances which may vitiate the results. Udransky<sup>2</sup> attributes the formation of coloured and resinous products in amyl alcohol to the presence of furfural, from which the alcohol can be purified, but the process is troublesome. The solvent action of amyl alcohol on urea and extractives

<sup>1</sup> *Rivista Italiana*, 1888

<sup>2</sup> *Zeitschr f physiol Chemie*, 1889

constitutes a further objection. Wormley<sup>1</sup> found that although amyl alcohol is nearly insoluble in water, 100 volumes of it agitated with water measure 109 volumes after the liquids have separated, he also found that amyl alcohol dissolves a certain amount of the *salts* of morphine from aqueous solution, the acetate more freely than the sulphate or the hydrochloride, but that the amount is diminished if the alcohol is previously saturated with water. As the result of a number of experiments, in which morphine was successfully extracted from urine with hot amyl alcohol, Wormley states that the presence of urea constitutes an almost insuperable difficulty as regards purification of the alkaloid. Acetic ether is not free from objections, it is soluble in water to a considerable extent (1 in 10), and dissolves extractives freely. The most convenient solvent consists of a mixture of equal parts of acetic and ethylic ethers, well washed by being shaken with water. Alkalisiation with sodium bicarbonate (excess of which does not redissolve morphine) should be effected in the presence of the solvent, and extraction accomplished without delay, if time is allowed for the precipitated morphine to become crystalline, it is much more resistant to the action of all solvents. Consult Tickle's<sup>2</sup> paper on the extraction of morphine by meta cresol.

**Tests.** A drop of strong nitric acid added to a fragment of morphine produces an orange-red colour. If a little of the alkaloid is dissolved in concentrated sulphuric acid and allowed to stand for fifteen to eighteen hours, and then treated with nitric acid a blue-violet colour, which changes to blood-red, is produced (Husemann). Sulphomolybdic acid (see test for brucine) produces a reddish-purple, which changes to blue. This and the previous test are the most delicate and conclusive for morphine, they will react to 0.1 milligramme. It is important to note that the *initial* colour-change is the one that is distinctive, the subsequent changes are common to several alkaloids. If a fragment of morphine is mixed with a couple of drops of strong sulphuric acid no coloration, or but the faintest pink is produced, the addition of a little ammonium selenate develops a pale yellow which changes to light green, sap-green, and brown. A small crystal of potassium bichromate, added to morphine dissolved in a drop of strong sulphuric acid, yields a green colour. If a little iodic acid is dissolved in a cubic centimetre of water in a test-tube, and an equal volume of carbon bisulphide is added, agitation produces in it no change of colour, the addition of a drop or two of solution containing morphine liberates iodine from the iodic acid, and on gentle agitation the carbon bisulphide, dissolving the free iodine, is tinged pink or rose-red. A drop or two of a solution of ferric chloride added to a strong solution of a salt of morphine produces a blue colour, if the reagent is in excess, the colour will be green, with morphine meconate this test produces a dark red—the reaction of meconic acid, which displaces the blue of the morphine. Reichard<sup>3</sup> mentions another reaction for morphine. On adding a trace of the alkaloid to a little titanous acid ( $\text{TiO}_2$ ) dissolved in concentrated sulphuric acid, an intense blackish-brown coloration is produced which becomes reddish-brown on agitation.

**Meconic Acid** may be tested for when the presence of opium, in which it exists in combination with morphine, is suspected. As just stated, it gives a red colour with ferric chloride, the colour is not destroyed by mercuric chloride, with lead acetate it gives a white precipitate which is soluble in nitric acid.

**Elimination** of morphine to a great extent takes place by the bowels, to a lesser extent by the kidneys, some experiments made by Alt<sup>4</sup> at the instigation of Hitzig show, in a striking way, the part played by the stomach in the process of elimination. A subcutaneous injection of morphine was given to a dog, in about four minutes after, the animal vomited, and the vomited matter was found to contain morphine. Further experiments show that, soon after morphine is subcutaneously introduced into the system, its elimination is

<sup>1</sup> *The Chemical News*, 1890

<sup>2</sup> *Pharm. Journal*, 1907

<sup>3</sup> *Zeitschr. f. anal. Chem.*, 1903

<sup>4</sup> *Berliner klin. Wochenschr.*, 1889

commenced by the gastric mucous membrane even when the stomach is empty and is continued until at least half of the dose injected is thus removed from the circulation, and is eventually voided with the *fæces*, a portion of the alkaloid is said to be excreted in the bile. Bougers<sup>1</sup> found that brucine, veratrine, caffeine, quinine, antipyrin, salicylic acid and other substances, when administered either subcutaneously or by the bowel, are eliminated by the stomach. In relation to such experiments Nencki<sup>2</sup> shows, from observations made on a dog which had a gastric fistula, that unless they are conducted with pure gastric juice an erroneous conclusion may be drawn, because, if bile be present, the reaction of the substance sought for might be obtained although the substance was not excreted by the stomach. In some experiments of the author's, conducted with the excretions from patients taking large medicinal doses of morphine, the alkaloid could always be detected in the *fæces* and occasionally in the urine, it is exceedingly difficult, however, to obtain morphine in the crystalline form, either from the tissues or the excretions of those who have been subjected to its influence. Morphine is excreted in the milk of women during lactation. Rosenthal<sup>3</sup> finds that morphine is eliminated in the saliva, and also that to some extent it may accumulate in the system.

The question of the decomposition of morphine in the organism must be regarded as undetermined, it is probable that some may be changed into oximorphine, or other derivative or combination of the alkaloid. For a chronological account of investigations on the subject, see Tauber<sup>4</sup> "*Ueber das Schicksal des Morphins im thierischen Organismus*"

### Chronic Poisoning by Opium and Morphine.

The habitual use of opium in gradually increasing doses produces an extraordinary degree of tolerance towards it, the practice of this habit is known as **opium-eating** or **morphinism**. According to Faust<sup>5</sup> the acquisition of this habit is not due to blunting of the tissues towards the action of morphine, but to a gradually increasing power of the tissues to destroy the drug. Thus, he found 70 per cent. of the early doses in the *fæces*, subsequently the amount became less, and eventually none at all was found. The morphine habit is frequently originated by the legitimate use of opium to relieve pain, and is afterwards continued on account of the agreeable sensations it produces, and to banish the feeling of depression which is experienced as soon as the effects of the drug pass off. A morphine *habitué* quickly **loses his moral perception**, and will descend to the lowest level of unblushing deceit in order to satisfy his cravings—when fully under the influence of the habit his moral fibre is disintegrated in all directions, ethically he is a coward, and avoids all methodical effort, if taxed with taking opium or morphine, he will deny the imputation with an earnestness and apparent sincerity that would convince anyone unacquainted with the facts. **Physical symptoms** develop after a time—visceral neuroses are evinced by the recurrence of violent pains in the region of the stomach and abdomen, which might be mistaken for those due to the passage of gall-stones, attacks of vomiting resembling gastric crises occur, and the bowels may be relaxed from time to time. The appetite is capricious and defective, considerable—sometimes excessive—emaciation takes place, and the patient looks shrunken and pallid in the face, like one suffering from

<sup>1</sup> *Arch. f. exp. Path.*, 1895

<sup>2</sup> *Arch. f. exp. Path.*, 1895

<sup>3</sup> *Centralbl. f. klin. Med.*, 1893

<sup>4</sup> *Arch. f. exp. Path. u. Pharm.*, 1890

<sup>5</sup> *Ibid.*, 1901

malignant disease, the later symptoms resemble those due to peripheral neuritis paræsthesiæ, neuralgia, trembling of the hands and ataxic gait, with numbness of the feet and tips of the fingers

The introduction of the hypodermic syringe has placed a convenient mode of self-administration of morphine within the reach of those disposed to make use of it. When the rapidity with which the system is brought under the influence of morphine injected subcutaneously is taken into consideration, the enormous doses that can be tolerated without producing the usual toxic effects are very astonishing. Stuart<sup>1</sup> records a case in which forty grains of morphine acetate were injected daily for months, even this amount has been exceeded.

The best **treatment** is to cut off the supply at once. This demands either great resolution on the part of the patient, or the bringing to bear of very powerful moral control on the part of a second person (which is seldom available), or, lastly, physical restraint. Conditions may exist which render gradual deprivation necessary, but the process is tedious and uncertain, when there appears to be danger of collapse from sudden total deprivation withdrawal of the drug in from six to twelve days may be resorted to. Unless there is some very cogent reason for doing so, it is better not to carry out "treatment" by substituting some other drug for morphine. For a considerable time after ceasing to take morphine the patient feels an intense craving for it, and he also suffers from severe gastric pain and dyspepsia, this is due to the fact that morphine is excreted by the stomach whether taken by the mouth or hypodermically, and consequently the sensory nerve-endings of the gastric mucous membrane, no longer benumbed by the constant presence of the drug, are hypersensitive to the gastric juice, which also tends to become abnormally acid, to counteract this the alkaline carbonates may be given. During and after the cure the patient must be restrained from the abuse of alcohol, otherwise an interchange of habits may occur.

## HEROIN.

Diacetyl morphine or heroin, a derivative of morphine, has, on several occasions, even when administered in proper doses, given rise to amaurosis, persistent vomiting and other dangerous symptoms. In a case recorded by Soles<sup>2</sup> about two and a half grains (the maximum dose being one-sixth of a grain) were accidentally administered, causing great prostration, diminished visual power with narrowly contracted pupils, slowing of the pulse, lowering of the temperature to 95° F, along with spasm and twitching of the limbs. Three one-and-a-half grain doses of caffeine citrate were injected subcutaneously and recovery took place.

Boyd<sup>3</sup> reports the following fatal case of poisoning by heroin --

A man, aged fifty nine, after eating a pie, had severe diarrhoea, and in three days' time became delirious. He was given 20 grains of potassium bromide, and an hour later, at 1.30 p.m., a cachet believed to contain barbiton, but which actually contained 6.9 grains of heroin. Two hours later he was tremulous, the face was twitching, and the skin acting freely. At 5 p.m. he was sleeping soundly, but the arms and legs were frequently jerked with strong clonic spasms. At 9.30 respiration became laboured and stertorous, the pupils

<sup>1</sup> *Brit. Med. Journ.*, 1889.

<sup>2</sup> *Allg. med. Centr. Zeitung*, 1899.

<sup>3</sup> *Medical Journal of Australia*, 1919.

were contracted to pin-points, and he could not be roused. The muscles of the back of the neck were rigid. At 6 p.m. the temperature was 101° F, pulse 100, respiration 14. The temperature gradually rose to 104° F before death, broncho pneumonia developed, and he died 70 hours after taking the heroin. The post mortem examination showed consolidation at the base of the lungs. The liver was in a condition of atrophic cirrhosis. The diarrhoea was probably due to food poisoning from the pie. This case illustrates the fact that heroin is more tetanising and less narcotic than morphine.

**Dionin**, or ethyl-morphine chloride, another derivative of morphine, has produced prolonged reddening of the face, a rash on the body closely resembling rubecula, and other toxic manifestations.

### BELLADONNA AND ATROPINE.

**Atropa Belladonna** (natural order *Solanaceæ*), or the deadly nightshade, contains the alkaloid atropine, to which it owes its toxic effects. Poisoning by belladonna occurs either from the improper use of one of the medicinal preparations of the drug, or from eating the berries in the fresh state.

**Atropine** ( $C_{17}H_{27}NO_3$ ) is a colourless crystalline substance with a strong alkaline reaction. It is odourless, sparingly soluble in water, but more so than most alkaloids, it is much more soluble in ether, still more so in spirit, and most of all in chloroform. The mydriatic alkaloids **atropine**, **daturine** (*datura stramonium*), **hyoscyamine**, and **hyoscyne** (*hyoscyamus niger*), **duboisine** (*duboisia myoporoides*) are all isomeric and probably convertible. Atropine can be split up into tropic acid and tropine, the latter being capable of forming combinations with other acids.

For the physiological action of atropine see p. 337. Elimination rapidly takes place by the kidneys. Dragendorff states that atropine will probably be found only in the urine which is first excreted after the reception of the poison.

**Symptoms.** The essential symptoms respectively produced by the plant and the alkaloid are identical, but when portions of the plant—as the berries—are eaten, nausea, vomiting and other signs of gastric irritation may be superadded. Belladonna poisoning is characterised by speedy onset of symptoms, which rapidly increase in severity. A hot, dry sensation, along with a feeling of constriction, is experienced in the throat, accompanied by thirst, the saliva is inspissated, and the tongue dry, swallowing is difficult or impossible. The pupils are dilated, usually to the uttermost, leaving only a narrow ring of the iris visible, they are insensitive to light, the conjunctivæ are suffused. The pulse, accelerated to 120 or 160 in the minute, is small and sometimes scarcely to be felt at the wrist. The skin, often covered with a scarlatina-like rash, which may be followed by desquamation of the epiderm, is hot and dry in the earlier stage, but may become cold during the stage of collapse. Alternate flushing and pallor of the face have been observed. In the early stage the patient complains of dizziness, indistinctness of vision—sometimes of diplopia; later on there may be complete loss of vision, he is unable to walk, and reels or stumbles on attempting to do so. Active delirium comes on, which often partakes of the imitative type, the patient will industriously perform a series of movements resembling those of a person employed in sewing with needle and thread, or he will tear off strips from an imaginary piece of cloth. In a case of poisoning by belladonna berries the patient imitated the acts of plucking fruit from a tree, conveying it to his mouth, and then swallowing it. These imitative acts are sometimes performed with such gravity and persistency

as to excite the risibility even of anxious friends who witness them. Hallucinations of sight are common, as may be gathered from the description of the mimic actions, sometimes the patient in the attempt to escape an imaginary danger will try to jump out of the window, or to rush through the door. The voice is stammering and the utterances are incoherent, but the patient is often extremely loquacious, intermitting his loquacity with wild peals of laughter, or vociferous shouting. Twitching of the muscles of the face and limbs often occurs, and may go on to tonic or clonic spasms affecting the entire body, which to some degree appear to be reflex in character, in a case recorded by Oliver<sup>1</sup> they were aggravated by the passage of the stomach-tube and by the withdrawal of the contents of the stomach. Sensory disturbances, such as numbness of the fingers, may be present. The urinary bladder and the intestines are usually paralysed.

In severe cases total insensibility with cold surface may supervene and last for many hours, on awakening, delirium has been known to recur. If the case is about to be fatal the insensibility increases, there may be recurrent convulsions or simply progressively deepening coma, death from paralysis of the heart and lungs occurring in from six to twenty-four or more hours.

When recovery takes place it is slow, several days elapse before the whole of the symptoms subside. The pupils continue dilated and only very gradually return to their normal size, accommodation may be defective for some time, and the memory is often enfeebled for three or four days, an aphasic condition existing in the meanwhile. A large percentage of cases of belladonna poisoning recover, even amongst those in which the symptoms are severe.

**Fatal Dose.**—One drachm of belladonna liniment, which was swallowed, and the same amount of the extract have respectively proved fatal. Recovery has taken place after a tablespoonful of the liniment and after ten drachms, also after half an ounce of the extract mixed with glycerine. Fourteen belladonna berries caused the death of an old man, recovery has taken place after fifty. Children are relatively less susceptible to the toxic action of belladonna than adults, recovery has followed in children after thirteen berries and even after thirty. In a case recorded by Strachan<sup>2</sup> a boy aged five years recovered after swallowing two drachms of extract of belladonna, five hours elapsing before treatment was commenced, in the course of which two quarter-grain injections of morphine sulphate were administered. An infant four months old recovered after the administration of half of a belladonna suppository, which would represent three-quarters of a grain of alcoholic extract of belladonna. Severe symptoms ensued, recovery being furthered by hot baths, stimulants, and minute doses of compound tincture of camphor (McWalter)<sup>3</sup>. Belladonna applied externally as a plaster or in the form of extract has often produced toxic effects. Howarth<sup>4</sup> records the case of a man who began to feel his mouth dry three-quarters of an hour after putting a belladonna plaster, six inches by four, on his back, the pupils were widely dilated, and delirium of the characteristic type occurred.

Two grains of atropine have proved fatal, severe symptoms have been caused by a quarter of a grain. In one case recovery followed three and a half grains of atropine sulphate, and in another five and a half grains. A child two and a half years old recovered from one quarter of a grain of atropine. Alarming and typical symptoms of poisoning have not unfrequently occurred after the instillation of a solution of atropine into the eyes for the purpose of

<sup>1</sup> *The Lancet*, 1891

<sup>2</sup> *Ibid.*, 1901

<sup>3</sup> *The Lancet*, 1903

<sup>4</sup> *Ibid.*, 1894

dilating the pupils. Homatropine, which is supposed to have none of the poisonous properties of atropine, is not free from risk. Hotz<sup>1</sup> states that the instillation of two drops into one eye, and three into the other, of a two per cent solution of homatropine rapidly produced scarlet-red colour of the face, headache, vomiting, and excitement. The following day the respirations were only five in the minute, the pulse was from 120 to 150. There was also extreme mydriasis. Recovery slowly took place under treatment. Brown<sup>2</sup> records pallor, dizziness, syncope, and slight delirium, the result of a solution of homatropine (0.6-300) having been twice dropped into the eye. Mydriasis lasted five days. Death has been caused by the application of atropine ointment to a blistered surface.

Although atropine is not often used criminally, a few cases of homicidal poisoning have occurred. One took place near Manchester, and was the subject of judicial inquiry at the Assizes held there in 1872 (*Reg. v. Steele*). The resident surgeon at a workhouse died from atropine poisoning, the alkaloid being detected in his body. He became ill after breakfast with typical symptoms and died in about twelve hours, the poison had been added to some milk used at breakfast which produced toxic symptoms in two other people who tasted it, a nurse, who it was alleged had a strong motive and the opportunity for committing the crime, was charged with the murder, but was acquitted. A curious case of belladonna poisoning is recorded by Bachner<sup>3</sup>. A man became ill after eating some soup prepared by his wife. When seen by a doctor he was flushed in the face, the eyes were bright and slightly bloodshot, the pupils were dilated and insensitive to light, the tongue was dry and covered with sticky saliva, the speech was stammering, the fingers trembling, the hands swollen, and the limbs cold. The patient complained of dizziness, singing in the ears, heaviness and heat in the head, indistinct vision, thirst, vomiting, and partial retention of urine. He recovered, and on judicial investigation it was found that he himself had added some belladonna seeds to the soup in order to accuse his wife of attempting to poison him.

**Treatment.**—When the poison had been swallowed the stomach-tube should be used and the stomach well washed out, if the instrument is not at hand an emetic must be given, after which stimulants and hot coffee are useful. Strong, stewed tea, or an infusion of tannic acid helps to precipitate and render innocuous any of the poison which may remain in the stomach. Douching of the burning, dry surface in the early stage, and artificial respiration in the stage of collapse are to be resorted to. Hypodermic injections of pilocarpine (one-third or half a grain of the nitrate or hydrochloride) tend to slow the pulse, tranquillise the respirations, and relieve the spasms if they are present. In default of pilocarpine a hypodermic injection of morphine (one-third, or half a grain) may be given. Binz<sup>4</sup> directs attention to the tolerance of morphine exhibited by cases of atropine poisoning, and instances this as being in favour of the antagonism of the two poisons.

**Post-mortem Appearances.**—In the absence of fragments of some parts of the plant there is no characteristic appearance in belladonna poisoning. If the berries have been eaten there may be signs of reddening of the mucous membrane of the stomach, the seeds should be carefully sought for in the stomach and intestines. The blood has sometimes been recorded as dark and fluid, with hyperæmia of the cerebral vessels, but such signs are of little value, the pupils usually remain dilated after death.

<sup>1</sup> *The Ophth. Rec.*, 1905

<sup>2</sup> *Ann. of Ophth.*, St. Louis, 1906

<sup>3</sup> *Friedreich's Blätter f. ger. Med.*, 1887

<sup>4</sup> *Centralbl. f. klin. Med.*, 1893



**Chemical Analysis**—Any seeds or fragments of leaves found in the stomach should be examined under the microscope. The seeds are small, ovoid or kidney shaped, and are covered with small projections which, under a low power, present a honeycombed appearance. The fresh berries are blackish purple in colour, and their juice stains a white surface purple, the mucous membrane of the stomach has sometimes been found thus stained. The addition of an alkali changes the purple to green, acids change it to red.

Atropine may be extracted from organic matter by the usual process, like morphine, it is soluble in excess of potassium and sodium hydroxide, it is very prone to undergo hydrolysis, especially in the presence of free alkalies. The evaporation of a solution containing atropine should be conducted at a temperature not exceeding 35° C, and excess of acid avoided. A mixture of three volumes of ether and one of chloroform is the best solvent for extracting the alkaloid from an aqueous solution.

**Tests.**—The chemical tests for atropine do not of themselves afford conclusive proof of the presence of the alkaloid, they give place to the physiological test, to which they are only to be regarded as corroborative. Alone among the ordinary fixed alkaloids free atropine reddens phenolphthalein, a minute fragment placed on phenolphthalein paper and moistened with a drop of water causes the paper to become red, if alcohol is dropped on the stain, the colour disappears, but returns when the alcohol has evaporated, the reddening of phenolphthalein paper by alkalies is not affected by alcohol. With a little atropine mix two or three drops of a strong nitric acid and evaporate to dryness over a water-bath, add to the yellow-coloured residue a few drops of alcoholic solution of potash—a reddish-violet or purple colour is produced. A fragment of atropine treated with sodium nitrate and strong sulphuric acid yields a yellow colour, which changes on the addition of alcoholic potash to reddish-violet, fading to pale rose.

**Physiological Test.** After proving the presence of an alkaloid by means of one of the alkaloid group reagents, a drop or two of a neutral aqueous solution prepared from the ether-chloroform extract should be instilled into the eye of a cat, or preferably a kitten, if atropine is present in only infinitesimal amount—0.01 mgrm—the pupil is dilated in from a few minutes to an hour, according to the amount present. The test may be repeated on the eye of a human being. Some of the urine voided during life or obtained from the bladder after death, if dropped into the eye of a kitten will cause dilatation of the pupil, thus affording a ready means of confirming the diagnosis. Cocaine also dilates the pupils when dropped into the eye, but a much stronger solution is required, and anaesthesia is also produced. The above reactions, chemical and physiological, are yielded by all the tropines.

Dragendorff detected atropine after it had been mixed with organic matter, which was then allowed to remain in a warm room two and a half months, until it was quite putrid. Ottolenghi<sup>1</sup> states that if atropine is exposed to the action of saprophytic bacteria it loses its mydriatic properties in four or five days.

## HENBANE.

**Hyoscyamus Niger**, or henbane, contains two basic substances—hyoscyamine and hyoscyne—which are isomeric with atropine. The fresh plant has a disagreeable odour, its juice, when dropped into the eye, dilates the pupils.

**Hyoscyamine** ( $C_{17}H_{23}NO_3$ ), with which **hyoscyne** is isomeric, is obtained from several atropaceous plants, it is convertible into atropine. Hyoscyamine is a colourless crystalline substance, without odour, moderately soluble in water, and freely soluble in spirit, ether, and chloroform, it has an alkaline reaction, and combines with acids to form salts.

<sup>1</sup> *Vierteiljahrsschr f ger Med*, 1896

**Symptoms.**—In many respects the symptoms produced by hyoscyamus are identical with those of belladonna, certain differences, however, have been observed. The face is flushed, the surface is hot and dry, the mouth and throat are parched, the pupils are enlarged and insensitive to light, vision is impaired, the pulse is quick and small, the respirations are of a sighing character, and in the early stage there is delirium. It has been noticed in cases of poisoning by hyoscyamine that the tendency to busy, wild delirium is not so great as with atropine. Trismus and clonic spasms of the muscles of the jaw and limbs have been observed. In the later stages the patient is comatose and collapsed, there is greater tendency to sleep and insensibility with hyoscyamine, and more especially with hyoscyne, than with atropine. Recovery is slow, as is the case with belladonna poisoning.

The **fatal dose** of henbane is not known, recovery has taken place after six drachms of the tincture were swallowed. Severe toxic symptoms followed the administration of one-tenth of a grain of hyoscyamine, and death after one-eighth of a grain swallowed along with the same quantity of morphine sulphate. A hypodermic injection of one-thirtieth of a grain of hyoscyne in one case, and the swallowing of about one-fortieth of a grain in another, produced severe symptoms of poisoning, followed by recovery. Even the maximum pharmacopœial dose may produce toxic symptoms. Worrall<sup>1</sup> records a case in which one-hundredth of a grain caused faintness, widely dilated pupils, palpitation, weak, rapid pulse scarcely to be felt and cold clammy skin, the condition seemed desperate. Four injections of pilocarpine (one-tenth of a grain in each) were administered, and after ten hours recovery took place. Given<sup>2</sup> records the case of a man, aged sixty-nine, who by a mistake of the dispenser, swallowed  $\frac{1}{16}$  of a grain of hyoscyne hydrobromide, instead of  $\frac{1}{60}$  as prescribed. In half an hour the patient was profoundly comatose, with stertorous breathing, the pulse (80) was small, the conjunctival reflex was abolished, and there was some twitching of the legs and arms. The stomach was washed out, strychnine and subsequently morphine were injected subcutaneously, and brandy and black coffee into the stomach and rectum. In eleven hours, consciousness returned, and complete recovery followed. In the case of *R v Crippen* (v p 69), Willcox estimated that there was in the organs submitted for analysis hyoscyne corresponding to two-fifths of a grain of hyoscyne hydrobromide. This would represent more than half a grain in the whole body. Some non-fatal cases of poisoning occurred through the accidental admixture of henbane seeds with celery seeds, for which they were purchased.

**Treatment.**—As in poisoning with belladonna.

The **post-mortem appearances** are not characteristic.

## STRAMONIUM.

**Datura Stramonium**, or thorn-apple, is another solanaceous plant which yields an alkaloid or alkaloids, that act much in the same way as those of belladonna and henbane. All parts of the plant are poisonous. The seeds are dark-coloured and kidney-shaped. They are about one-eighth of an inch in length, and have a rough surface.

**Daturine** is an isomer of atropine. According to Ladenburg the kind known as "light daturine" consists principally of hyoscyamine.

**Symptoms.**—The following case related by Steiner<sup>3</sup> illustrates the toxic

<sup>1</sup> *Australasian Med Gaz*, 1889

<sup>2</sup> *The Lancet*, 1904

<sup>3</sup> *Berliner Klin. Wochenschr.* 1887

action of stramonium. A man aged forty-five drank a decoction prepared by boiling the leaves and fruit of the *Datura stramonium*, to relieve pain in his chest. About three-quarters of an hour afterwards he sprang out of bed and ran about the room, looking into all the corners like a person bereft of reason, he was put to bed by force and held there as he made violent efforts to escape, being all the while unconscious. The face was red, the pupils were widely dilated and did not react to light, the limbs moved spasmodically, the pulse, which intermitted, was 130 in the minute, the respirations were deep and quickened, the skin was dry, and the temperature 96.6° F. Swallowing was difficult, cutaneous sensibility was abolished, the abdomen was distended but not painful on pressure. The patient subsequently lay quiet and became comatose, the redness of the face changing to pallor, the respirations became quieter and slower, and the pulse sank to 120. The patient appeared to be dying, but shortly after began to improve and gradually recovered, weakness and trembling of the limbs persisting for a week. A man swallowed a teaspoonful of Himrod's asthma-specific, which is intended for inhalation, many of the symptoms of stramonium poisoning occurred, but the pulse was very slow, only twenty-five to the minute, recovery took place. The remedy contains stramonium with probably lobelia and potassium nitrate.

The **fatal dose** is not known. About one hundred seeds, and seventeen or eighteen grains of the extract, have caused death. Death has occurred in seven and in twenty-four hours.

**Treatment** As in poisoning with belladonna.

The **post-mortem appearances** are not characteristic.

**Chemical Analysis**—After isolation and proof of the presence of an alkaloid, its physiological effect should be tested as with atropine, the same chemical tests may also be used.

The tiopines, respectively derived from belladonna, hyoscyamine, and stramonium, being isomeric, and possessing the same chemical properties, identification of the group, one of which was the poison administered, is all that can be accomplished in a toxicological investigation, there is no reliable test by which any one of these alkaloids, in minute quantity, can be distinguished from another.

**Duboisine**, obtained from the leaves of the *Duboisia myoporoides*, is another isomer of atropine. It is considered by some to be identical with hyoscyamine, Ladenburg believes it to be identical with hyoscyne. It is a powerful mydriatic, and produces symptoms like those of atropine.

A case is recorded by Chadwick<sup>1</sup> in which one-hundredth of a grain of duboisine sulphate, dropped into the eyes of an old man, produced dizziness, weakness, and loss of control of the legs, dryness, with bitter taste in the mouth, huskiness and indistinctness of speech, visual hallucinations—the patient grasping at imaginary objects in the air, and glancing suspiciously under the bed-clothes and behind his back—slow pulse, and a copious flow of words without relevancy. Kollock<sup>2</sup> relates an almost parallel case caused by the instillation of two drops of a solution of duboisine sulphate (one grain to two drachms) into the eyes. The face was flushed, the pupils dilated, the patient was dizzy and moved himself from side to side, though apparently rational he made remarks devoid of meaning and relevancy, and was unconscious of all that had occurred whilst in this condition.

The **treatment** is the same as for atropine.

<sup>1</sup> *Brit. Med. Journ.* 1887.

<sup>2</sup> *Med. News* 1887.

## SOLANUM.

**Solanum Nigrum**, or garden nightshade, and **Solanum Dulcamara**, or bitter sweet, contain an alkaloid—solanine, the latter also contains dulcamarine.

**Solanine** is an irritant, and when swallowed, causes gastro enteritis, it also acts on the nervous system, though somewhat irregularly, probably owing to the presence in plants, from which it is derived, of varying amounts of tropines of unknown composition, which may be responsible for some of the atropine like symptoms which occasionally occur.

Poisoning by substances containing solanine occurs from eating the berries of the bitter-sweet or other plants of the same genus. Vomiting and diarrhoea, with more or less collapse, pain in the stomach, cramps in the legs, followed by clonic spasms, dilatation of the pupils, pallor, coldness of the surface, hallucinations, and coma are amongst the symptoms which may be met with. The respiratory function is lowered, and in fatal cases the respiratory centres are paralysed, death taking place from asphyxia.

The **treatment** consists in furthering the evacuation of the stomach—vomiting almost invariably occurring spontaneously—the administration of stimulants, and possibly opium, with the application of warmth.

**Solanum Tuberosum**, or the common potato, has, on several occasions, acted as a poison. Cortial<sup>1</sup> records an instance in which 101 soldiers were affected, and Schmiedeberg<sup>2</sup> four similar instances occurring in garrisons, in one of which 357, in a second 90, in a third 125, and in a fourth 43 men were thus poisoned. The symptoms comprised—frontal headache, colicky pains in the stomach and bowels, vomiting, diarrhoea, tenderness of the abdomen, shivering, sweating, depression, slight stupor, oral and visual disturbances, congestion followed by pallor of the face, blueness of the lips, widely dilated pupils, initial acceleration with subsequent retardation of the pulse, elevation of temperature, syncope, and convulsions. All the 716 men thus attacked recovered. The potatoes which acted in this untoward way presented no abnormal appearance. Banks<sup>3</sup> records an instance in which four members of a family, after eating a large quantity of potatoes, were simultaneously attacked with pain in the abdomen and back, difficulty in micturition, shivering, cold surface, and swelling of the abdomen, which was tender on pressure, all recovered. In this case the potatoes were unsound, three additional members of the family, who also ate freely of them, but who took the precaution of previously removing the diseased portions, were not affected. Morris<sup>4</sup> records the case of a girl of fourteen who ate some berries of the potato plant, and a few hours after began to have pain in the abdomen, she subsequently became collapsed, with hurried respiration and quick feeble pulse, the pupils were not dilated, death took place on the third day.

The symptoms of poisoning in these cases were possibly due to the presence in the potatoes of an unwanted amount of solanine. Meyer<sup>5</sup> analysed a number of ordinary potatoes, and also some taken from the same store as those which poisoned the 357 men as recorded by Schmiedeberg, with the following results—Good potatoes contain about 0.044 grm. of solanine per kilogramme of the unpeeled tuber, some young potatoes contained as much as 0.236 grm., and some newly budded from seed potatoes contained no less than 0.580 grm. per kilo. Potato sprouts, and young green potatoes are rich in solanine.

Savage<sup>6</sup> has recently reviewed the records of poisoning attributed to potatoes, and does not consider the view that the symptoms were due to solanine poisoning is completely justified, or at least is proved, evidence pointing more to the potatoes having been infected with bacteria. He describes an outbreak in which 80 to 100 persons who had eaten potatoes and fish tried in oil developed symptoms. The potatoes were good, since half of them bought the evening before were eaten without harm. The remaining half were washed and scraped and left ready for frying on the following day. In this unprotected state they were probably infected in some unascertained way. This inoculated nutrient medium was kept for at least 16 to 20 hours in a hot shop in August, and was then fried with fish. Probably the cooking in the oil killed the bacteria and left their toxins.

The **treatment** is symptomatic. The bowels should be freed from the undigested potatoes, of which there is often a large amount, and the gastro intestinal symptoms then relieved with opium.

<sup>1</sup> *Arch. de Med. et de Pharm. Militaire*, 1887.

<sup>2</sup> *Arch. f. exp. Path. u. Pharm.*, 1895.

<sup>3</sup> *Dublin Quarterly Journ. of Med. Sc.*, 1846.

<sup>4</sup> *Brit. Med. Journ.*, 1859.

<sup>5</sup> *Arch. f. exp. Path. u. Pharm.*, 1895.

<sup>6</sup> *Food Poisoning and Food Infections*, 1920.

## INDIAN HEMP.

**Cannabis Indica**, or Indian Hemp, is a deliriant and hypnotic, and has been used in the form of *hashish* to procure sensuous hallucinations

**Cannabin**, an active principle prepared from it, is a brown syrupy liquid which has the odour of Indian hemp **Cannabinon** is a dark brown resin which has sedative properties

The **symptoms** of an overdose are thus described by a medical man who took forty drops of the tincture Giddiness and fulness in the head, heaviness and numbness of the feet and legs, complete loss of sensation as far as the knees, which rendered standing difficult and walking impossible, the same symptoms commenced in the tips of the fingers and reached to the elbows, but the anæsthesia was not so complete as in the legs Anxiety and dread of death were experienced, and the heart's action was tumultuous and irregular The mental condition was emotional laughing and crying alternating, no pleasurable excitement was experienced In a case recorded by Casaccia<sup>1</sup> two drachms of the alcoholic extract produced the following symptoms in half an hour – Mental exaltation with a tendency to physical movements, paræsthesiæ of hands and feet, heat in the epigastrium, dryness of the fauces dilatation of the pupils, which reacted to light, and full, slow pulse of fifty-eight to the minute The patient talked incoherently without intermission uttering cries or howls at intervals, recovery took place

The **fatal dose** is not known Seven and a half minims of the tincture have caused toxic symptoms Death has occurred in twelve hours, it may be delayed for several days, in one case it did not occur until the nineteenth day Un-toward effects have also been observed after the medicinal use of Cannabinon

The **treatment** is the same as for opium

## GELSEMIUM.

**Gelsemium Sempervirens**, or the yellow jasmine of North America, owes its toxic properties to the presence of an alkaloid **gelsemine**, which when instilled into the eye is a powerful mydriatic, when administered internally in small doses it contracts the pupils, in poisonous doses it dilates them It paralyses the spinal cord and the respiratory centres, and produces tetanus specially affecting the facial muscles and the muscles of articulation, it also gives rise to ataxic symptoms, it has little effect on the heart and brain Gelsemine is eliminated by the kidneys

**Symptoms.**– A case in which a comparatively small quantity of the tincture produced toxic symptoms is related by Jepson<sup>2</sup> A woman who had been previously taking the drug without obtaining any benefit took an increased dose of twenty minims of tincture of gelsemium every three hours for three or four doses She lost power over her tongue, being unable either to articulate or to swallow except with great difficulty, the pupils were widely dilated and vision was indistinct, she had a sensation of uncertainty in the movements of her hands and arms, but retained consciousness, and recovered under treatment Myrtle<sup>3</sup> prescribed some pills containing one-tenth of a grain of *gelsemin* (the powdered alcoholic extract of gelsemium root, the dose of which is from half a grain to two grains), for which the dispenser substituted hydrochloride of the alkaloid *gelsemine*, the dose of which is from a sixtieth to a twentieth of

<sup>1</sup> *Riv di Chim Med e Farm*, 1883

<sup>2</sup> *Brit Med Journ*, 1891

<sup>3</sup> *Brit Med Journ*, 1889

a grain The patient became giddy, was sick, and lost the power of speech, the tongue was drawn to one side, the muscles on the right side of the face quivered, and she could not guide her hand, trismus, clonic spasms, exhaustion, and unconsciousness for two hours were amongst the symptoms, recovery took place Three teaspoonfuls of the fluid extract of gelsemium caused the death of a woman in seven and a half hours

**Treatment.**—If taken by the mouth the poison should be evacuated, either by the tube or an emetic, then stimulants should be administered, warmth applied, and artificial respiration resorted to if necessary Atropine and strychnine have been recommended as antidotes in order to stimulate the respiratory centre

**Chemical Analysis**—Separation from organic matter is effected as with the alkaloids in general, ether or benzene may be used to shake out from aqueous solution

**Test**—Gelsemine has a bitter taste If a fragment of the alkaloid is dissolved in a few drops of strong sulphuric acid no coloration is produced a granule or two of manganese dioxide stirred into the mixture develops a deep crimson-red which changes to green

## COCAINE.

**Cocaine** ( $C_{17}H_{21}NO_4$ ), benzoyl methyl-ecgonine, one of several alkaloids yielded by the *Erythroxylon coca*, is a colourless crystalline substance, it has a bitter taste and leaves a sensation of numbness on the tongue It is only slightly soluble in water, much more soluble in alcohol, and still more so in ether, benzene, and chloroform

It is largely used as a local anæsthetic, and acts in that capacity by paralyzing the terminals of the sensory nerves, it blanches mucous membranes, and produces some dilatation of the pupils

Taken internally cocaine first stimulates and then paralyzes the nerve-centres of brain and cord With poisonous doses the action of the heart, in animals, is slowed and the blood pressure reduced, the respiratory function after an initial increase is lowered and finally paralysed The temperature is raised, and convulsions may occur Although it has been found in the urine, cocaine probably undergoes decomposition in the body into ecgonine free or combined

**Symptoms.**—The following case related by Haenel<sup>1</sup> illustrates the course of acute cocaine poisoning A dentist injected into the gums of a girl of nineteen a solution equal to about one gram and a third of a salt of cocaine, to lull the pain of tooth-extraction The patient became pale and then fell down and was severely convulsed, she was unconscious, the pupils were widely dilated and reactionless to light At first the pulse was too quick to be counted, subsequently it dropped to 176 in the minute, the temperature was 100.8° F, and the respirations were 44 in the minute The patient remained unconscious for seven hours, and on regaining consciousness experienced diminished sensibility of the hands anæsthesia of the mucous membrane of the mouth and nostrils, with loss of taste and smell, during the first twenty-four hours there was retention of urine The stimulating effects of the poison on the respiratory centres with paralysis of the vagi would account for the heart and lung disturbance Walker<sup>2</sup> saw a man, aged twenty-four, about four and a half hours after he had accidentally swallowed between eight and nine grains of cocaine

<sup>1</sup> *Berliner klin Wochenschr*, 1888

<sup>2</sup> *The Lancet*, 1895

hydrochloride The man appeared as though partly under the influence of alcohol, he complained of a sense of constriction in the throat and over the heart, and of a painful numbness in the stomach and abdomen with an indefinite feeling of oppression and mental dulness The pupils were dilated, and did not react to light, the pulse was uncountable Forced muscular movements of the limbs occurred, and the body was rotated from side to side and bent at the same time, involuntary movements of the muscles of mastication caused the patient to present the appearance of a person chewing tobacco The first urine passed after taking the poison was of a green colour There was occasional dyspnoea with lividity of the lips, which was relieved by amyl nitrite, a strong purgative produced some liquid tarry motions, and the following day the patient appeared quite well Garland<sup>1</sup> states that a girl of seventeen almost immediately after swallowing from twelve to fifteen grains of cocaine in solution, experienced vertigo followed by nine rapidly succeeding epileptiform convulsions, death taking place within forty minutes Montalti<sup>2</sup> records the case of a woman who took twenty-three grains of cocaine hydrochloride, fifteen minutes after delirium came on, she tried to vomit, but did not succeed, rigors occurred, the face was pale, the pupils were dilated and the lips cyanotic, she became pulseless and unconscious and died forthwith Zambianchi<sup>3</sup> states that a woman had about three and a half grains of cocaine injected into the breast preparatory to an operation, she immediately had epileptoid convulsions and died in twenty minutes Palmer<sup>4</sup> states that in less than an hour after a man, aged forty, had swallowed ten grains of cocaine hydrochloride, the eyeballs were protruded and immobile, and the respirations sank to eight in the minute, recovery took place

**Fatal Dose.**—About two-thirds of a grain injected subcutaneously caused the death of a woman aged seventy-one in five hours Curgenvén<sup>5</sup> records the case of a woman who, after swallowing ten grains of cocaine hydrochloride in solution, became cyanosed the pulse being quick and the respirations shallow, tetanic spasms occurred, and she died in one, between forty and fifty minutes after the poison was swallowed On the other hand, a man habituated to the use of cocaine, injected under his skin twenty-three grains daily for some time In another case recovery took place after forty-six grains were taken into the stomach Death has occurred from the injection of a solution of cocaine into the tunica vaginalis The injection of one drachm of a 4 per cent solution of cocaine hydrochloride into the urethra caused immediate dilatation of the pupils, flushing, twitching of the face, and convulsions, death occurred four minutes after the first convulsion In another case related by Mathieson,<sup>6</sup> twenty minims of a 4 per cent solution injected into the urethra caused immediate convulsions and death The action of cocaine is very irregular Weinrich<sup>7</sup> relates two cases in which all but fatal results, including Cheyne-Stokes respiration, followed the injection into the urethra of a solution containing two grammes of cocaine, although, in one case, the same amount had been previously injected on six separate occasions without any unusual effect These two were the only cases out of several thousand, similarly treated, in which any toxic effects were produced

**Treatment.**—If the poison was swallowed, the stomach must be evacuated and washed out with the aid of the tube, if it was injected under the skin or

<sup>1</sup> *Ibid*, 1895

<sup>2</sup> *Lo Sperimentale*, 1888

<sup>3</sup> *Gazz degli Ospidali*, 1888

<sup>4</sup> *The Lancet*, 1898

<sup>5</sup> *Quarterly Med Journ*, 1896

<sup>6</sup> *Dublin Journ Med Sc*, 1895

<sup>7</sup> *Berliner klin Wochenschr*, 1896

beneath a mucous membrane, treatment is limited to the administration of stimulants, with the inhalation of chloroform, if necessary, to relieve the spasms which interfere with respiration. Artificial respiration may be required.

**Post-mortem Appearances.**—The principal changes—which are due to vaso-motor paralysis—are hyperæmia of the membranes of the brain and cord, and of the viscera generally.

**Chronic Poisoning** by cocaine occurs in those who have acquired the habit of injecting the alkaloid hypodermically, in the same way in which morphine *habitues* use morphine, also when long used as a spray to the nasal and laryngeal tracts, in one case it was due to the repeated application of vaginal tampons containing cocaine. In recent years the habit of snuffing cocaine powder has become very common among certain classes in Paris. A number of ill-effects are produced on both the moral and the physical well-being of those who have become victims to the habit, mental apathy and moral degeneration are accompanied by disturbances of the digestive organs, anomalous pains, and general emaciation. Cutaneous anæsthesia of the extremities may occur, numbness and tremulousness of the fingers producing a certain amount of clumsiness and want of co-ordination. Anæsthesia of the pharynx may give rise to a sensation as of a foreign body. The speech may be jerky and explosive. As stated by Savage,<sup>1</sup> the cocaine-taker develops hallucinations of his senses which are readily detected and which call attention to his vice, he hears whispers and voices, whilst the morphine-taker sees things. A symptom of chronic cocaine poisoning, known as Magnan's symptoms, is produced by disturbance of sensation—the patient complains of feeling as though grains of sand, or small round bodies, or in some instances worms, were under the skin. The will power is lost, he becomes irritable and quarrelsome, and may learn to indulge in alcohol to excess.

**Tests.** A solution of a salt of cocaine has a bitter taste, and causes loss of sensation in the tongue and other parts of the mouth with which it comes in contact. Mezger<sup>2</sup> recommends the following test.—To a solution of cocaine hydrochloride in water add a few drops of a 5 per cent solution of chromic acid, as each drop of the chromic acid solution is added a precipitate is formed which immediately redissolves, if a small quantity of strong hydrochloric acid is now added a heavy yellow permanent precipitate is formed. Several alkaloids are precipitated from neutral solution by chromic acid—strychnine, brucine, veratrine, quinine, for example—but no alkaloid except cocaine requires the addition of hydrochloric acid after the chromic acid before permanent precipitation takes place. If a little cocaine is treated with a few drops of strong nitric acid, and the mixture is evaporated to dryness on a water-bath on adding a few drops of a strong alcoholic solution of soda or potash to the residue and stirring them well together, an agreeable, aromatic, ethereal odour is given off somewhat resembling that of the flower called the meadowsweet. Ammonia throws down a flocculent white precipitate from a not too weak solution of cocaine, the precipitate is soluble in excess, but in a short time the solution deposits long needle-shaped crystals, which agglomerate in feathery groups. If a little potassium chromate dissolved in strong sulphuric acid be added to the residue obtained by evaporating to dryness a solution of cocaine, a red colour is produced which changes to green, this, on the addition of a few drops of water, becomes greenish yellow. A drop of solution of ferric chloride, added to a solution of cocaine, yields a faint yellow, which becomes orange colour or red on boiling. If a moderately strong solution of cocaine be heated with

<sup>1</sup> *The Lancet*, 1905

<sup>2</sup> *Chem. Centralblatt*, 1890.



half its volume of sulphuric acid, an odour of benzoic acid is evolved, when cold, if the solution be shaken out with ether, and the ether be separated and evaporated, crystals of benzoic acid are left behind

### MALE FERN.

**Felix Mas**, or male fern, is extensively used as an anthelmintic in cases of tapeworm. It contains filicic acid, an amorphous, white, tasteless powder without smell, which is probably the active principle of the rhizome. Experiments on animals made by Poullsson<sup>1</sup> show that filicic acid produces tetanic convulsions, followed by paralysis, the convulsions resembling those produced by strychnine, heart paralysis occurs along with the general paralysis, although the heart may beat a few times after respiration has ceased.

**Symptoms.**—A man, thirty years old, was given a draught containing one ounce and a half instead of one drachm and a half of the extract of male fern, which he took in two doses. Soon after the first dose he felt unwell, and after the second, which was given some hours subsequently, he began to vomit, and was purged, then followed cramps, profuse sweating, delirium, and coma, which ended in death about twenty hours after the draught was taken. At the necropsy the omentum and the peritoneal covering of the small intestines were bright red, and in the submucous tissue of the stomach were ecchymoses with linear extravasations on the surface of the mucous membrane. Meyer<sup>2</sup> reports the case of a man twenty-eight years of age, who, after a "moderate dose" of extract of male fern, became comatose, and remained so for a day and a half. When he recovered consciousness he was quite blind of the right eye and nearly so of the left, due to optic neuritis. Atrophy of both optic nerves ensued. Stulp<sup>3</sup> also reports coma, followed by snow-white oedema of the fundus, which resulted in optic atrophy. Freyer<sup>4</sup> relates an instructive case in which a child, aged two and three-quarters years, took eight capsules—each containing about fifteen grains of extract of male fern, along with the same quantity of castor oil—in five hours, she became somnolent, and as though paralysed, and died after the occurrence of some spasms. Section showed petechial ecchymoses in the mucous membrane of the stomach, pronounced injection of the mucous membrane of the intestines, and venous filling of the various organs. The interesting point to note is that three weeks previously the child took double the quantity of the extract, *but without the castor oil*. A case is recorded by Hofmann,<sup>5</sup> in which a child, five and a half years old, had very nearly two drachms of the extract given to her in three draughts, death took place in six hours, with symptoms of trismus and general spasms, much the same appearances were found as in the other cases.

The case recorded by Freyer has a practical bearing. The toxic properties of the extract of male fern are augmented by the presence of additional oil to that contained in the extract itself, the same child tolerated twice as much of the extract alone as that which proved fatal when given in combination with castor oil, it is advisable, therefore, not only to avoid giving a mixture of the extract with castor oil, but also to give some other laxative than oil, if one is subsequently needed. A case is recorded by Schlier<sup>6</sup> in which an adult very nearly lost her life, probably owing to a tablespoonful of castor oil being

<sup>1</sup> *Arch. f. exp. Path.*, 1891.

<sup>2</sup> *Deutsch. med. Zeitung*, 1905.

<sup>3</sup> *Zeitschr. f. Ther. u. Hyg. d. Augen*, 1904.

<sup>4</sup> *Therapeutische Monatshefte*, 1889.

<sup>5</sup> *Wiener klin. Wochenschr.*, 1890.

<sup>6</sup> *Munchener med. Wochenschr.*, 1890.

given one hour after a draught which consisted of extract of male fern mixed with the powdered root

**Treatment.**—If spontaneous vomiting does not occur, the stomach should be emptied either by the tube or an emetic, after which general treatment will be required, and probably the administration of stimulants

### LOBELIA.

**Lobelia Inflata**, or Indian tobacco, contains a basic substance, *lobeline*, which is the active principle of the plant. Lobeline is an oily, yellowish-coloured fluid with a burning taste, it is soluble in ether, and slightly so in water, it resembles nicotine in many of its properties

When taken in large doses lobelia acts as a depressing emetic, like tobacco. Cases of poisoning by lobelia chiefly result from its free administration by quacks. In a case recorded by Wharton and Stillé,<sup>1</sup> a woman was poisoned by a quack giving her half a teacupful of infusion of lobelia, seeds and all, she died in half an hour, and on examination the stomach was found to contain a table spoonful of lobelia seeds, its mucous membrane was softened and much inflamed, the intestines were also inflamed. In another case<sup>2</sup> one drachm of the powdered leaves was given by a quack, great pain was produced, with vomiting, small pulse, contracted pupils, insensibility, spasmodic twitchings of the face, collapse, and death in thirty-six hours, in this case also the mucous membrane of the stomach was found much inflamed

The **treatment** consists in emptying the stomach in those exceptional cases in which vomiting has not occurred spontaneously, and then administering stimulants freely. Warm applications should be made to the surface, and the recumbent posture maintained until the heart has quite recovered itself

**Chemical Analysis**—The basic principle may be extracted with ether from an alkaline aqueous solution

**Tests.**—On evaporation of the ether, the residue gives a violet coloration with sulphomolybdic acid, this is like the reaction of morphine, but the fluidity, odour, and colour of lobeline will prevent any confusion between the two, moreover lobeline turns red on the addition of strong sulphuric acid—a reagent which does not affect morphine

### TOBACCO.

**Nicotiana Tabacum**, or tobacco, contains an alkaloid nicotine—in combination with malic or citric acids—upon which its toxic properties depend

**Nicotine** ( $C_{10}H_{14}N_2$ ), liberated by the action of alkalis from tobacco, is closely allied to pyridine, it is a colourless, volatile, oily liquid which turns brown and resinous on exposure to air. It has a marked alkaline reaction, and forms salts with acids, it is freely soluble in water, alcohol, and ether, and has an acrid taste, and a strong odour like that of the juice of an old well-used pipe

After first stimulating the vagus both centrally and peripherally, thus slowing the heart-beats, nicotine paralyses the cardiac terminals and causes rapid, irregular action of the heart. The respiratory rate is first accelerated and then retarded. The peripheral blood-vessels are contracted by poisonous doses of nicotine, hence the pallor and coldness of the surface. Nicotine first

<sup>1</sup> *A Treatise on Med Jurisprudence*, 1860

<sup>2</sup> *Pharmaceutical Times*, 1874

stimulates and then paralyzes the cerebral and spinal centres. With small doses the pupils may at first be contracted, but they are dilated when the toxic symptoms are fully developed. Nicotine is to some extent eliminated in the urine.

The **symptoms of acute poisoning** by swallowing tobacco juice or nicotine are — A burning, acrid sensation in the throat, a sudden feeling of depression, with giddiness, loss of power over the limbs, faintness, nausea, vomiting, tremors, coldness of the surface with clammy sweat, loss of consciousness, contraction of the pupils, which in fatal cases are subsequently dilated, feeble irregular action of the heart, laboured sighing respirations, complete relaxation of the whole musculature, with possibly delirium and convulsions. Before the patient loses consciousness he may experience a feeling of oppression or of sinking in the cardiac region, accompanied by great anxiety, dimness of vision, and loss of power of speech. Occasionally the bowels and the bladder are involuntarily evacuated. In some instances the lethal action of the poison is exceedingly rapid, in one case death occurred in eighteen minutes, in another in three or four minutes. In the well-known case of Fougner, who was poisoned in 1850 by his brother-in-law, Count Bocarmé, with nicotine which he prepared for the purpose, death took place in five minutes.

The leaves of the tobacco plant applied to the unbroken skin have caused symptoms of poisoning, an infusion of them similarly applied in order to kill parasites, has on several occasions caused death. The infusion injected into the rectum as a vermifuge, has frequently proved fatal, on one occasion after only twelve drops, and on another after an infusion prepared from half a drachm of tobacco, an infusion of about 35 grains of green tobacco leaf in half a litre of water caused the death of a six-year-old girl in half an hour. Even smoking tobacco has caused acute fatal poisoning, although most of the nicotine present is converted into pyridine-bases during the combustion of the tobacco. A boy smoked a pennyworth of twist tobacco and afterwards became very sick and fell in the street, he went home to bed and at four o'clock in the morning vomited again, three hours after he was found lying on the bed, dead and cold<sup>1</sup>. Two or three drops of nicotine taken into the stomach would probably be fatal in a few minutes. A drunken man was killed by his companions emptying the juice of their pipes into some spirit and giving it him to drink. Tobacco in the form of infusion or juice has caused death in from twenty minutes to seven or eight hours.

**Treatment.**—If the poison was swallowed the stomach-tube should be used, or an emetic given, followed by stimulants, external warmth, artificial respiration if necessary, and the maintenance of the recumbent posture. Hypodermic injections of strychnine (one twenty-fifth grain) have proved serviceable. Strong tea, or a solution of ten or twenty grains of tannin in water may be given.

**Post-mortem Appearances.**—The odour of tobacco is usually perceptible on opening the abdomen. When the poison has been swallowed the mucous membrane of the stomach may be injected or ecchymosed, the intestines have been found contracted and to contain blood-stained mucus.

**Chemical Analysis.**—Nicotine may be separated from organic admixture by the usual process for the isolation of alkaloids. Ether is the best solvent after its evaporation the residue consists of oily looking drops.

**Tests.**—Nicotine is freely soluble in water. If a solution of mercuric chloride is added to an aqueous solution of nicotine a white precipitate is formed which

<sup>1</sup> *The Lancet*, 1885

subsequently becomes yellow and crystalline. Silver nitrate produces a white precipitate which subsequently becomes black. Chlorine-water added to an aqueous solution of nicotine produces no turbidity. A little ethereal solution of iodine added to an ethereal solution of nicotine produces an oily mass in which red crystals are formed, which have a watch-spring lustre when viewed by reflected light. If a trace of nicotine is mixed with a drop of a 40 per cent solution of formic aldehyde, a solid deposit is formed after some hours which gives an intense rose coloration when touched with a few drops of nitric acid (Schindlmeister).<sup>1</sup> The odour of nicotine and its toxic effects on animals afford further means of identification.

**Chronic nicotine poisoning** results from excessive smoking, and also from the inhalation of tobacco-charged air in manufactories. The symptoms are dyspepsia, anæmia, and nervous disorders, among which amblyopia, with contraction of the field of vision and with central scotomata for red and green, intermittent tumultuous action of the heart, and a tendency to faintness and dizziness are the most prominent. Bury<sup>2</sup> has seen three cases of peripheral neuritis due to excessive smoking of tobacco.

### SPOTTED HEMLOCK.

**Conium Maculatum**, or spotted hemlock, so named from dark purple spots on the stem, is a plant belonging to the natural order *Umbelliferae*, its leaves resemble parsley sufficiently to cause them to have been mistaken and eaten for that plant. It has a peculiar, very characteristic "mousy" odour, which can be developed by pulpifying the leaves or other part of the plant in the presence of a little caustic soda or potash. The plant contains two alkaloids—conine and methyl-conine—along with other bases.

**Conine** ( $C_8H_{17}N$ ) is a colourless oily liquid, which turns brown on exposure to air, it possesses the "mousy" odour of the plant in a high degree, and has an acid bitter taste. It is strongly alkaline and combines with acids to form salts, it is sparingly soluble in water, and is freely soluble in alcohol, ether and chloroform.

Conine paralyzes the motor nerve-terminals, and subsequently the motor centres of the brain and spinal cord, the paralysis spreading from the periphery to the centre. Death is due to respiratory paralysis, and is usually preceded by asphyxic convulsions. Conine is eliminated in the urine.

**Methyl-conine** ( $C_9H_{19}N$ ) abolishes the reflex of the spinal cord.

**Symptoms.**—A burning sensation with a feeling of constriction in the throat is experienced, followed by nausea, vomiting, pain, oppression in the stomach and bowels, and diarrhoea. The nerve-symptoms are variable, probably on account of differences in the relative proportion of conine and methyl-conine present in the poison swallowed. Progressively increasing muscular weakness, with dyspnoea, the respiratory movements becoming slower and slower, without disturbance of the higher centres, are the symptoms usually met with, but sometimes delirium, coma, and partial convulsions are prominent from the first. The pupils are dilated, and the surface of the body is cold. In the pure motor-paralysis type the patient first feels weakness in the legs which causes him to stumble when trying to walk, this deepens into complete paralysis which creeps up towards the trunk, the arms being less rapidly affected. The paralysis eventually invades the muscles of respiration, the patient becomes cyanotic, and death takes place from dyspnoea.

<sup>1</sup> *Rev. intern. Falsif.*, 1901.

<sup>2</sup> *The Lancet*, 1896.

In the pure paralytic type convulsions are not infrequent during the final stage, but they are due to asphyxia caused by the respiratory paralysis. The sensory nerves are relatively but slightly affected.

A peculiar case is recorded by Schulz<sup>1</sup> of a student who, after repeatedly smelling at some conine, experienced weakness of the limbs, inability to keep the eyes open, burning sensation of the conjunctivæ, pain in the head, affection of speech, general feeling of heat followed by profuse perspiration, he rambled and was unable to sleep. The headache continued for twenty-four hours, along with a tendency to perspire profusely on the least movement. Gunn<sup>2</sup> records the case of a woman who, after inhaling for four or five minutes the vapour given off from a drachm of tincture of conium in boiling water, complained that her limbs felt heavy and powerless, the pupils were dilated and vision was impaired, urgent dyspnoea with cyanosis necessitated artificial respiration. The pulse and the intellect were unaffected, for some hours after there was a tendency to recurrence of the dyspnoea.

**Treatment.**—Evacuate the stomach and then give stimulants and apply warmth. Artificial respiration is sure to be required in severe cases, and should be persistently kept up, life may be saved by this means when the condition appears almost hopeless.

**Post-mortem Appearances.**—In the absence of traces of the poison in the viscera, there is no characteristic appearance. The blood will probably be dark and fluid, with other indications of death from asphyxia.

A child, eight months old, had given to it one teaspoonful of a mixture containing one drachm of extract (prescribed in mistake for succus) of conium with one drachm of potassium bromide in an ounce and a half of chloroform water. When seen the legs were paralysed, occasional twitchings of the arms and head occurred, but no decided convulsions, the pupils were dilated, the face was livid and the respirations were diaphragmatic, death took place in seven hours. Pepper, who made the post-mortem examination, found the organs generally congested, along with an increased amount of serum in the cerebral ventricles and under the arachnoid, and an injected condition of the membranes of the spinal cord. The right heart was distended with blood, the bases of the lungs were hyperæmic, and punctiform extravasations were observed on the surface of the liver. The contents of the stomach yielded no odour of conium until they were treated with potassium hydroxide and heated, when the "mousy" odour became apparent. An ethereal extract obtained from the contents of the stomach, when treated with hydrochloric acid, yielded crystals of conine hydrochloride.<sup>3</sup>

**Chemical Analysis.**—Separation of conine from organic admixture may be accomplished as with nicotine. Considerable caution is necessary in the identification of conine, since substances somewhat resembling it may be obtained from cadavers that have undergone putrefactive changes, any such products, however, do not yield the chemical reactions of conine, nor are they strongly toxic, they probably consist of or contain cadaverin—a ptomaine which has an odour somewhat resembling that of conine, but is scarcely so "mousy."

**Tests.**—Conine is less soluble in hot water than in cold, therefore if a cold saturated aqueous solution is heated it becomes turbid, like albuminous urine, similarly treated, but, unlike the urine, it clears up again on cooling. If conine is exposed to the vapour of hydrochloric acid, crystals of conine hydrochloride are formed. A few drops of a solution of mercuric chloride added to a solution

<sup>1</sup> *Deutsche med. Wochenschr.*, 1887.

<sup>2</sup> *The Lancet*, 1894.

<sup>3</sup> *The Lancet*, 1885.

of conine in water produce a white amorphous precipitate, which does not change to yellow nor become crystalline as is the case with the precipitate formed by nicotine when similarly treated. Silver nitrate produces a dark-brown precipitate, which turns black. Chlorine water added to an aqueous solution of conine produces turbidity. Treated with chromic acid, conine yields butyric acid, which may be recognised by its odour.

### CENANTHE CROCATA.

**Cenante Crocata**, or water dropwort, is another umbelliferous plant, of which the leaves and especially the root possess powerful toxic properties. Pohl<sup>1</sup> obtained from the root a substance—*œnanthotoxin*—which is soluble in alcohol, ether, chloroform, and all the usual solvents except petroleum ether, it is insoluble in water, in dilute alkaline solutions and in acids. A rabbit weighing 830 grms. died in convulsions half an hour after receiving 0.02 gm. of *œnanthotoxin*.

The symptoms comprise convulsions, cyanosis, insensibility, laboured respirations, collapse, dilated pupils, delirium, small, feeble, slow pulse with gastro enteric disturbance, on one or two occasions the convulsions have been of a strychnine like character. In some instances the symptoms have been almost entirely psychical, consisting of hallucinations, wild laughter, and actions like those seen in delirium tremens. Death may occur very quickly in one case a man died in five minutes, and in another in a quarter of an hour after the symptoms commenced. Two other cases died on the ninth and eleventh day respectively with gastro enteric symptoms. Cows and horses are also poisoned by it. A waggoner ate some *œnanthe* to cure scurvy, his horse eating some at the same time, the man died in an hour and a half, and the horse in two hours and a half.

### CICUTA VIROSA.

**Cicuta Virosa** or water hemlock, is a poisonous umbelliferous plant which, like *Cenante crocata*, has been eaten in mistake for parsnip and for celery. It produces symptoms like those produced by *Cenante crocata*. From it Pohl separated a substance—*cicutotoxin*—which possesses similar toxic properties to *œnanthotoxin*. Boehm<sup>2</sup> states that in some respects *cicutotoxin* produces effects analogous to those produced by strychnine and picrotoxin.

### FOXGLOVE.

**Digitalis Purpurea**, or foxglove, is a plant belonging to the natural order *Scrophulariaceæ*, the leaves of which possess toxic properties due to the presence of three glucosides—digitalin, digitalein, and digitonin—with one other active principle. A variety of preparations are sold under the name of **digitalin**, which differ in chemical constitution and physiological effects, in accordance with the mode in which they are obtained. The most poisonous of the active principles is digitoxin, which is not a glucoside.

Digitalis is essentially a heart poison, and causes death from heart paralysis, the pulse usually ceasing before respiration, the respiratory rate is often slowed, especially when death is imminent. The active principles of digitalis probably undergo decomposition in the body, very exceptionally traces of them have been found in the urine.

**Symptoms.**—Primarily the digestive tract is affected by a poisonous dose either of digitalis or of its active principles. Nausea, vomiting—which is often very obstinate and persistent—pain, with a sensation of oppression in the region of the stomach, thirst, and colicky pains in the abdomen, with or without diarrhoea, are common. After a varying interval the more specific effects of the poison manifest themselves—giddiness, with a feeling of faintness, headache, increased oppression in the epigastric region, moisture and coldness of the

<sup>1</sup> *Arch. f. exp. Path.*, 1894.

<sup>2</sup> *Ibid.*, 1876.

surface especially of the limbs, prostration, and various affections of the special senses, as dimness of vision, noise in the ears, are present, with which mental disturbances, in the form of hallucinations or delirium, may be associated. The action of the heart is profoundly affected—the pulse sinks hour by hour in rapidity and tension, and becomes very intermittent and fluttering. The respirations are slow, and assume a sighing character. If the patient lifts his head when in the recumbent posture a tendency to syncope asserts itself, and if he stands upright actual syncope probably occurs, which may prove instantaneously fatal. An inclination to somnolence, which may deepen into coma, is not unfrequent, cyanosis, with or without asphyxic convulsions, may precede death.

It is to be noted that the special action of digitalis on the heart renders the patient liable to fatal syncope for several days after the immediate effects of the poison have passed off. In the acute stage, the pulse-rate may be lowered to under forty beats per minute, in the case of a woman who drank some infusion prepared from fresh digitalis leaves, the pulse sank to thirty-six, with periods of entire cessation of the heart's action at short intervals.

**Fatal Dose.**—Nine drachms of the tincture of digitalis have proved fatal, and recovery has taken place after more than three times as much. Thirty-eight grains of the powdered leaves have caused death, and recovery has followed one drachm. The fatal dose of digitalin is not known. Mawer<sup>1</sup> relates the case of a woman who swallowed fifty-six granules, each containing one milligramme of Homolle's digitalin, the total dose being equal to eighty-four grains of digitalis leaf, the effects produced were giddiness, vomiting, pain in the stomach, dusking of the face, dilatation of the pupils, coldness of the extremities, oppression in the præcordial region, slow respirations with prolonged inspiration, and slow, irregular, weak pulse, which sank to forty-four in the minute, recovery took place. Radcliffe<sup>2</sup> records the recovery of an infant, one year and eleven months old, after a milligramme and a quarter of Nativelle's digitalin. Death has taken place in twenty hours, but it may be delayed to a much more remote period.

**Treatment.**—If necessary use the stomach-pump or give an emetic, such as mustard or zinc sulphate with hot water. Stimulants should be freely given and external warmth applied, the patient being kept in the recumbent posture for several days. Hot applications, friction, or mustard leaves, to the epigastrium are useful. Hot coffee with brandy in it may be given. If the vomiting is prolonged ice in small quantities will be useful.

The **post-mortem appearances** are not characteristic. There may be some signs of irritation, or of inflammation of the gastric mucous membrane.

**Chemical Analysis.**—Fragments of the leaf, should the poison have been taken in that form, may be detected in the stomach, in which case they should be examined microscopically.

The aqueous extract obtained in the usual way from organic matter is best shaken out with chloroform, in which all the active principles of digitalis are soluble, they are not all soluble in ether nor in benzene, it is to be remembered that digitalin in acid solution is taken up by chloroform.

**Tests.**—If digitalin is dissolved in a little concentrated sulphuric acid, and some bromine-water is added to the mixture, a violet-red colour is produced. A little digitalin gently heated with a few drops of a mixture of equal parts of sulphuric acid and alcohol turns yellow-brown, on the addition of a drop of a dilute solution of ferric chloride a green or bluish-green colour is produced.

<sup>1</sup> *The Lancet*, 1880

<sup>2</sup> *Brit. Med. Journ.*, 1901

The physiological test may be resorted to as performed by Tardieu in the celebrated case of Pommereais, who was convicted of fatally poisoning a woman with digitalin. Three frogs were prepared so that their hearts were exposed — One frog was left unpoisoned, into the pleural sac of the second a solution of digitalin was injected, and into that of the third some of the suspected poison obtained from the body of the deceased, the heart-beats of the three frogs were respectively counted at stated intervals. The heart of the unpoisoned frog showed little change, that of the one to which digitalin was administered promptly and progressively slowed until it ceased to beat, the heart of the frog to which the suspected poison was administered behaved like number two, excepting that the effects were less rapidly produced.

**Strophanthus** —Müller<sup>1</sup> records that a patient with kidney disease, forty-seven years of age, swallowed from two to three drachms of tincture of strophanthus. Unconsciousness, tetanic and clonic convulsions, hallucinations, anæsthesia, myosis, diarrhoea, and Cheyne-Stokes breathing occurred, and death ensued on the fourth day. (The symptoms and course of this case are suggestive of uræmia.)

### COLCHICUM.

**Colchicum Autumnale**, or meadow saffron, is dependent for its toxic effects upon the presence of an active principle, colchicine, with a small trace of veratrine, both of which are chiefly contained in the root and the seeds.

**Colchicine** ( $C_{22}H_{25}NO_7$ ) is a yellowish crystalline powder when pure, but is often met with as an amorphous resinous-looking substance. It is soluble in water, and freely so in alcohol and chloroform, it is slightly, if at all, soluble in ether, and is insoluble in petroleum ether. Colchicine is decomposed by acids with the exception of tannic acid, with which it combines.

Colchicine in poisonous doses causes irritation of the nerve-endings in the intestines, along with gastro-enteritis. The motor-centres in the cord and medulla are paralysed, and death is produced by paralysis of the respiratory centres; the sensory nerves are also paralysed. From experiments on animals Jacobi<sup>2</sup> concludes that colchicine may be converted into oxydicolchicine within the organism. Colchicine is partly eliminated by the kidneys and bowels, chiefly by the latter.

**Symptoms.**—A burning pain in the throat which extends down the œsophagus to the stomach, where it assumes an aggravated form, is experienced shortly after the poison is swallowed, then follow copious vomiting and purging, the latter being accompanied by violent colicky pains in the abdomen. There is intense thirst, the face is shrunken and pallid, or cyanosed, the surface is cold and moist, the pulse is small, irregular and rapid, the breathing is slow and laboured—the whole symptoms in fact resemble those due to an attack of cholera, the resemblance being increased by the nature of the evacuations from the bowels, which, after the normal contents are discharged, chiefly consist of serous fluid, subsequently they become blood-stained. A sensation of oppression is felt in the region of the heart, the patient is profoundly depressed, and, being fully conscious, suffers greatly. Muscular twitchings or spasms may occur, the whole body occasionally being convulsed, the pupils are sometimes dilated, sometimes contracted, strangury may be present, with increased or diminished amount of urine. Towards the end the cyanosis often becomes more marked, the collapse then being very profound, the mind usually remains clear till towards the last, in exceptional cases stupor occurs earlier.

<sup>1</sup> *Inaug. Dissert.*, Berlin, 1898.

<sup>2</sup> *Arch. j. exp. Path. u. Pharm.*, 1890.



**Fatal Dose.**—Three and a half drachms of colchicum wine have caused death. Recovery has taken place after ten drachms, which produced severe toxic symptoms. The lethal dose of colchicine is not known, a woman, aged forty-three, swallowed about six grains, which had been substituted for another drug, and died in thirty-one hours (Albertoni e Casali<sup>1</sup>). Death has taken place in seven hours, it usually occurs within thirty hours, but has been delayed for three, and even seven days.

**Treatment.**—The stomach should be emptied by the tube, and well washed out with a solution of tannic acid, or an emetic may be given, followed by strong tea, then brandy, by the mouth, or if vomiting forbids, ether injections, external warmth and friction, with artificial respiration if required. Probably a subcutaneous injection of morphine will be advisable to relieve the intense colicky spasms of the bowels.

**Post-mortem Appearances.**—They are not characteristic, there may be signs of inflammation in the mucous membrane of the stomach and bowels, possibly with spots of ecchymosis, but in some cases there has been an entire absence of such indications.

**Chemical Analysis**—Advantage may be taken of the insolubility of colchicine, in petroleum ether, to dissolve out fatty substances from an aqueous solution obtained from the organic matter. Colchicine is dissolved out of acid solution by chloroform. The chloroform solution may either be evaporated to dryness, or after it has undergone some degree of concentration, petroleum ether may be added so as to cause the colchicine to crystallise out.

**Tests.**—A drop of nitric acid, sp gr 1.4, brought into contact with colchicine, produces a violet colour, which changes to brownish-yellow. One part of ammonium vanadate dissolved in two hundred parts of sulphuric acid produces a green coloration (sometimes very evanescent, and not distinct except with the pure alkaloid), which changes to a brownish-violet, the reagent should be freshly prepared. The physiological test does not afford decisive information, the conclusion arrived at by a committee of French experts who were appealed to in a case of suspected colchicine poisoning, was that experiments on animals do not afford the means of determining that poisoning by colchicine has taken place.

Ogier<sup>2</sup> was able to obtain the reactions of colchicine isolated by the usual process, from the exhumed bodies of dogs which he had poisoned with it five and a half months before. In the bodies of animals poisoned with it, Obolonski<sup>3</sup> detected colchicine four and a half months after death.

## VERATRUM.

**Veratrum Album**, or white hellebore and **Veratrum Viride**, or green hellebore, contain a number of alkaloids, Wright and Luff<sup>4</sup> found jervine, pseudo-jervine, rubi-jervine, cevadine, veratralbine, and veratrine. Commercial veratrine is an impure alkaloid obtained from sabadilla seeds.

**Veratrine** ( $C_{37}H_{53}NO_{11}$ ) is a white, crystalline powder, having an acid, burning taste, when it comes in contact with the nasal mucous membrane it excites violent sneezing. It is insoluble in water, and is soluble in ether, chloroform, and spirit. It has an alkaline reaction.

Veratrine first stimulates the motor nerves, and then paralyses their endings. It alters the character of muscular contractility, contraction is prolonged,

<sup>1</sup> *Bollet delle scienze med.*, 1890

<sup>2</sup> *Annales d'Hygiène*, 1886

<sup>3</sup> *Vierteljahrsschr f. ger. Med.*, 1888

<sup>4</sup> *Journ. Chem. Soc.*, 1879

and relaxation takes place slowly—a condition resembling, but not identical with, tetanic spasm. The sensory nerves also undergo primary stimulation, followed by paralysis, which is more complete than is the case with the motor endings. The activity of the heart is reduced, the vaso-motor apparatus paralysed, and the blood-pressure consequently lowered. Respiration is first quickened, then slowed, and finally arrested from paralysis of the respiratory centres, and probably also of the vagus endings in the lungs. The result of all this is that the temperature is lowered. Veratrine is quickly eliminated by the kidneys.

**Symptoms.**—An acrid, burning sensation, with constriction, is experienced in the throat, the burning sensation extends along the œsophagus down to the stomach, and is followed by vomiting and great thirst. Diarrhœa is not invariable, but may occur, if it does there is usually tenesmus. The pulse is feeble, and the respirations are slowed and sighing in character, the pupils are sometimes dilated. Pallor and coldness of the surface, with rapid collapse, twitching of the muscles, and even convulsions, have been observed. Giddiness and paræsthesiæ, followed by superficial anæsthesia, may occur in the early stage, consciousness is usually maintained until the stage of collapse is reached, but occasionally, early on, there is a tendency to delirium and stupor.

**Fatal Dose.**—Not known. In one case death took place after about eighteen grains of the powdered root of *V. album*, recovery has occurred after more than twelve times that amount. Grenander<sup>1</sup> records the case of a woman who drank some liniment containing four and a half grains of veratrine. The pupils were dilated, the pulse was slow (50 per minute) and feeble, the respirations were slow and shallow, consciousness was not impaired, salivation and profuse sweating occurred, vomiting was frequent, great oppression was felt in the epigastric region, together with soreness of the throat, and profound prostration, there was no diarrhœa, recovery took place under prompt treatment. In another case reported by Blake<sup>2</sup> nearly three grains of veratrine were accidentally swallowed by an adult. The patient complained of giddiness, sickness, constriction of the throat, thirst, diarrhœa with tenesmus, and a tired, weak, faint feeling, the tongue was swollen and the mouth and throat were sore, the pupils were extremely contracted, the respirations hurried, and the pulse was quick and small, micturition was frequent. A continued tingling was felt over the entire body, with now and then intolerable fits of itching in different parts, there was no sneezing. Recovery took place under treatment, the irritation of the skin being the last symptom to subside.

**Treatment.**—After evacuation of the stomach with the tube or an emetic, stimulants and hot coffee should be administered. External warmth and friction may be required, with maintenance of the recumbent posture and artificial respiration. If excessive diarrhœa is present, morphine will be advisable.

**Post-mortem appearances** are not characteristic, only few reports are extant, and they afford no definite information.

**Chemical Analysis.**—Chloroform, or a mixture of chloroform and ether, is the best solvent to extract veratrine from aqueous solution. It can be shaken out of an acid solution, but more perfectly after the addition of alkali.

**Tests.**—Applied to the mucous membrane of the nostrils veratrine causes violent sneezing. A drop or two of strong sulphuric acid added to a little veratrine in a watch-glass and well mixed, develops a yellow colour, which quickly changes to orange and finally to cherry-red, if the mixture is warmed.

<sup>1</sup> *Hygiea*, 1885

<sup>2</sup> *St George's Hosp Reps*, 1870

it becomes red immediately Salicine treated with sulphuric acid turns red immediately without heating Narcotine gives a similar reaction, but takes hours to acquire the red colour Hydrochloric acid with veratrine produces no change until the mixture is heated, when it becomes red Sulphomolybdic acid added to a fragment of veratrine produces a brick-red, which becomes dirty brown, greenish, and finally blue If a little veratrine is mixed with five or six times the amount of cane sugar, and moistened with concentrated sulphuric acid, a yellow colour is first produced, which changes to green and finally to blue With ammonium selenate and sulphuric acid veratrine yields a brownish-yellow colour which changes to rose-red

### MONK'S-HOOD.

**Aconitum Napellus**, or monk's-hood, sometimes called wolf's-bane, is a common plant belonging to the natural order *Ranunculaceæ*; it is extremely poisonous in all its parts The root has been eaten for horse-radish, although the difference between the two is so marked as to make it impossible for any observant person to mistake one for the other - Aconite-root quickly tapers to a point, whereas horse-radish is cylindrical or thereabouts, aconite-root is brown, horse-radish is a dirty white On section, aconite-root is soft in texture, and white in colour, the cut surface quickly changing to pink on exposure to air, horse-radish is tough and white, and retains its colour unchanged The taste also of the two roots is different aconite is acrid and imparts a tingling sensation, followed by numbness, to the tongue and lips, with a feeling of constriction in the throat, horse-radish is simply pungent

The aconite plants contain a number of alkaloids and derivatives, which have been investigated by Wright, Luff, and Menke,<sup>1</sup> several of them not being poisonous Commercial aconitines consist of variable admixtures of some of these alkaloids, and therefore greatly differ in potency, English and French aconitines are the strongest, German aconitine is much less powerful Recent investigations by Dunstan, Passmore, and Umney<sup>2</sup> indicate that aconitine is mono-benzoyl aconine An exhaustive account of the aconite bases is contained in Allen's *Commercial Organic Analysis*, vol III, part II, 1892 The physiological action of these bases has been experimentally investigated by Cash<sup>3</sup>

**Aconitine** ( $C_{33}H_{45}NO_{12}$ ) is one of the most active, if not the most active, poison known, it is crystallised with difficulty, and is usually met with in white amorphous masses, it has an alkaline reaction, and forms salts, of which the nitrate is preferred English aconitine is but slightly soluble in water, and is not very freely soluble in alcohol and ether, whilst the German alkaloid is soluble in all three, and freely so in ether

German aconitine has a bitter, sharp, burning taste, the English alkaloid is not bitter, but is sharp and burning All aconitines produce a peculiar tingling and numbness of the lips and tongue, which comes on shortly after a drop of a dilute solution is applied to them, the sensation lasts for some time, and is very characteristic of the poison

When introduced into the system in poisonous doses aconitine produces a general tingling all over the body, the parts liberally supplied with sensory nerves being most affected The poison first stimulates and then paralyses the sensory nerve-terminals, it produces the same effect on the motor nerves

<sup>1</sup> *Journ. Chem. Soc.*, 1877, 1879

<sup>2</sup> *Proc. of the Chem. Soc.*, 1892.

<sup>3</sup> *Philosophical Trans. Roy. Soc.*, 1898.

and centres of the medulla and cord, the higher cerebral centres are little affected. The heart-beats, at first retarded, may subsequently be quickened, the motor ganglia and the muscular substance of the heart are eventually paralysed. Respiration becomes slow, and afterwards shallow, due to the action of the poison on the respiratory centre. Death is generally due to arrest of respiration, after cessation of which the heart may continue to beat for a short time. The temperature sinks from the first. Aconitine is eliminated in the urine and probably in the feces, in experimenting with animals Diagen-dorff found it in both. Cash<sup>1</sup> states that the peripheral innervation of the respiratory muscles is not interfered with.

**Symptoms.**—Shortly after swallowing a poisonous dose of a preparation of aconite, tingling, followed by numbness of the lips, mouth, and throat, is experienced, due to the direct contact of the poison with the parts affected, then a feeling of nausea and pain in the stomach develops, which is usually followed by vomiting and sometimes by purging. A tingling, numb sensation—due to the poison which has been absorbed—is now felt over the whole body, with giddiness, imperfect vision, restlessness, anxiety, twitching of the muscles (sometimes with spastic contractions), darting pains in the legs, and muscular prostration. The pulse is feeble and intermittent, the respirations are laboured and spasmodic, and the temperature sinks, the limbs especially being cold and moist to the touch. The pupils may alternately dilate and contract, and there may be delirium, or a tendency to drowsiness and stupor, towards the end convulsions may occur, which are probably not altogether asphyxic. In an instance of multiple aconite poisoning related by Baker,<sup>2</sup> in which four boys, from fourteen to eighteen years of age, chewed pieces of aconite root the symptoms developed in from a few minutes to half an hour, all the patients felt heavy and sleepy, and experienced most of the symptoms just described, in the worst case the pupils were widely dilated, the respirations were spasmodic, but the pulse though small was quiet and regular, all recovered.

**Fatal Dose.**—One drachm of aconite root, two grains of the Pharmacopœial extract, and one drachm of the tincture have respectively proved fatal. The smallest recorded fatal dose was eighty minims of the Pharmacopœial tincture taken in ten doses, spread over four days, the largest individual dose being ten minims, this is quite an exceptional case. Twenty-five minims of Fleming's tincture, equal to about two drachms of the official tincture, have proved fatal, and recovery has followed one ounce. A fatal case of poisoning by aconite liniment is recorded by M'Whannel.<sup>3</sup> A woman swallowed one ounce of the Pharmacopœial liniment (equal to about five and one-third drachms of dried aconite root), and became collapsed, with small, irregular pulse, slow laboured breathing, cold clammy limbs, and pallid lips. There were no convulsions, the pupils dilated immediately before death, which took place in sixty-five minutes after the poison was swallowed. Death has occurred in from three-quarters of an hour up to fifteen or even twenty hours after the poison was swallowed, the usual period of survival being from three to four hours.

As regards the fatal dose of aconitine, experience is more limited. An instructive case is recorded by Tresling.<sup>4</sup> A medical man who had prescribed aconitine nitrate, was informed that the medicine produced strange symptoms, and, in order to demonstrate its harmlessness, took a dose himself equal to about one fifteenth of a grain, in about an hour and a half after, he began to feel ill. When seen four hours after he was pale, the surface was

<sup>1</sup> *Loc cit*

<sup>2</sup> *Brit Med Journ*, 1882.

<sup>3</sup> *Brit Med. Journ*, 1890.

<sup>4</sup> *Weekbl. van het Nederl. Tijdschr. v. Geneesk.*, 1880.

cold, the pupils were contracted, the pulse was small and irregular, but not rapid, the tongue was swollen, and the patient experienced a burning sensation in the throat, with pain down to the stomach, headache, weakness of the limbs, and shivering. The pupils suddenly dilated, and synchronously there was loss of vision, shortly after the pupils resumed their previous condition, vision returning. Vomiting occurred both spontaneously and in consequence of emetics. In four hours and forty minutes a convulsion occurred, followed by a succession of others, respiration became more laboured, and the pupils again dilated with accompanying loss of vision. Later, the vomiting became very violent, unconsciousness supervened, the pupil remained dilated and insensitive to light, the respirations grew slower, and the heart ceased to beat, death occurring five hours after the aconitine was taken. At the necropsy pallor of the skin and of the muscles was observed, with hyperæmia of the stomach and first part of the intestines, the colon was pale and the rectum bloodless, the lungs were hyperæmic, and the heart contained fluid blood, the cerebral membranes were injected, the ventricles contained blood stained serum, and blood was extravasated on the choroid plexus, the blood throughout was fluid and cherry red in colour. Death was attributed to heart paralysis. In this case French aconitine (Petit's) was dispensed in place of a weak German preparation (Friedlander's), by experiments on animals, Plugge<sup>1</sup> afterwards found that the alkaloid dispensed was one hundred and seventy times more potent than that which was prescribed.

The *cause célèbre* of aconite poisoning was that of *Reg v Lamson* (C C C, 1882), the prisoner being a medical practitioner who was accused of having poisoned his brother in law. He paid a visit to his victim, a boy of nineteen, who was a boarder in a school, and persuaded him to swallow a gelatine capsule, which he pretended to fill with sugar, but which, as the result showed, contained aconitine (Morson's). In about twenty minutes the boy complained of heartburn, and then vomited, he had great pain in the stomach, a sense of constriction in the throat, was restless, and tossed himself violently about whilst in bed, the breathing became slower, the heart's action feebler, and he died about four hours after swallowing the capsule. At the necropsy the membranes of the brain were slightly congested, but there was no fluid under them nor in the ventricles, the lips were pale, the pupils dilated, the lungs congested, especially at the lower part, the heart was empty, the liver, spleen, kidneys, and mucous membrane of the stomach and the first part of the duodenum were congested, on the surface of the gastric mucous membrane were six or eight small, slightly raised patches. From a portion of the vomited matter, from the urine obtained after death, and from the viscera, Stevenson and Dupré obtained aconitine, which responded to the usual physiological tests. The prisoner was condemned and executed.

A fatal case of poisoning due to aconite and belladonna combined is recorded by Lipscomb.<sup>2</sup> A girl of seventeen swallowed two tablespoonfuls of a liniment composed of equal parts of the aconite and belladonna liniments of the Pharmacopœia. The face and neck were flushed, the neck, arms, and to a slighter degree the legs were convulsed, the movements being aggravated by external stimuli, the pupils were dilated, the heart's action was quick, turbulent, and irregular—probably 300 per minute, the radial pulse could not be felt, and in one hour and forty minutes the heart suddenly ceased to beat, respiration continuing for a few seconds longer.

**Treatment.**—Evacuate the stomach with the tube or an emetic. Administer stimulants freely—brandy by the mouth or the rectum, ether subcutaneously. External warmth, friction, artificial respiration, and the recumbent posture will be required.

**Post-mortem Appearances.**—Not characteristic, see the accounts already given. If the poison was taken in the crude form, search should be made for fragments of the root or other parts of the plant. In a case in which six persons were poisoned, three fatally, by the accidental addition of aconitine to quinine-wine, the only special feature noticed at the necropsy was the presence, in all three, of subpleural ecchymoses.<sup>3</sup>

**Chemical Analysis.**—Separation from organic matter is accomplished by the usual process, in the course of which exceptional care is necessary to prevent decomposition of the alkaloid, which easily undergoes hydrolysis. The alcoholic extract is preferably made without the addition of an acid, in any case a mineral acid must not be used.

<sup>1</sup> *Arch der Pharm*, 1882.

<sup>2</sup> *Brit Med Journ*, 1888.

<sup>3</sup> *Annales d'Hygiène*, 1892.

**Tests.**—After proving the presence of an alkaloid, the only reliable procedure is to make use of the physiological test. A tingling sensation, followed by numbness of the lips or tongue, produced by the application to them of a drop of a solution of the product obtained from the vomit, excreta, or tissues, is strongly indicative of aconitine, the subsequent administration of a known quantity of the solution to one of the smaller animals, the toxic effects being compared with those produced by aconitine on other animals of the same species and weight, will yield sufficient evidence of its presence.

Opposite opinions are given with regard to the permanency of aconitine in the presence of putrefying organic matter. Lewin maintains that it is not destroyed. Stevenson states that it cannot be detected if allowed to remain some time along with decomposing animal matter which has become alkaline.

### HELLEBORE.

**Helleborus Niger**, or true hellebore, has a dark coloured root which is sometimes used as a vermifuge by herbalists and others, the leaves are also used for the same purpose. The toxic properties of hellebore depend upon the presence of two glucosidal active principles, **helleborin** and **helleborein**, both of which tend to produce muscular paralysis and to cause vomiting and diarrhoea. Helleborin acts on the brain and causes insensibility, it also produces local anaesthesia, and if applied to the nostrils occasions sneezing. Helleborein produces first slowing and then quickening of the heart, and also dyspnoea.

**Symptoms.**—A stinging, numb feeling of the tongue extending to the throat, with colicky pain in the stomach and abdomen, followed by violent vomiting and purging, are experienced, together with dizziness, heavy sensation in the head, drowsiness, prostration, collapse, with cold, pallid, perspiring surface, feeble pulse, and laboured respiration, in the event of a fatal issue, death may be preceded by convulsions, the pupils are frequently dilated. In a case recorded by Hlott,<sup>1</sup> a young man put about two teaspoonfuls of powdered hellebore into some water and drank it off. He was seized with violent cramps, giddiness, dimness of vision, inability to stand, and profuse vomiting, the pulse was only 40 in the minute, the pupils were dilated, a burning pain was felt in the epigastrium and a sensation of constriction in the throat, the fauces were red and swollen, recovery took place.

**The fatal dose** is not determined. Half a drachm of a watery extract is recorded as having been fatal. Death has occurred in from three to twelve hours.

**Treatment** consists in promoting evacuation of the stomach, followed by the administration of stimulants and morphine to allay excessive action of the bowels, external warmth should be promoted.

The **post-mortem appearances** are not characteristic, as with other vegetable irritants, signs of inflammation in the mucous membrane of the stomach have been observed.

**Chemical Analysis**—Helleborin, but not helleborein, may be shaken out of *acid* aqueous solution with ether, it is still more soluble in chloroform. After evaporation of the solvent, the residue immediately yields a bright red colour on being touched with a glass rod which has been dipped in strong sulphuric acid.

<sup>1</sup> *Brit Med Journ*, 1889

## STAVESACRE.

**Delphinium Staphisagria**, or stavesacre, a plant belonging to the natural order *Thalimifloræ*, yields seeds which contain several active principles, among which are the two alkaloids, **delphinine** and **staphisagrine**, in toxic action the former resembles aconitine and the latter curare

Poisoning with stavesacre is exceptionally rare. A case is recorded in which a man by mistake swallowed two teaspoonfuls of a powder, two thirds of which consisted of powdered stavesacre seeds. The heart was slowed to 35 or 40 beats per minute and was very feeble in its action, severe collapse came on, the surface being very cold, the breathing was laboured, the pupils were dilated, and the abdomen was distended and exceedingly painful, consciousness was undisturbed. Under treatment recovery took place in a few hours<sup>1</sup>

## LABURNUM.

**Cytisus Laburnum**, or common laburnum, contains an alkaloid—**cytisine** ( $C_{20}H_{27}N_3O$ ), which has basic properties and forms salts with acids. Cytisine is freely soluble in water, alcohol, acetic ether, and chloroform, but is insoluble in ether. It has a bitter taste, and is powerfully toxic. Cytisine first stimulates and then paralyzes the cord and motor nerves, the paralysis beginning in the peripheral endings, the respiratory centres also are first stimulated and then paralyzed, death being due to respiratory paralysis. The heart-beats are accelerated. After slight excitation of the brain, cytisine produces somnolence and coma. Cytisine is rapidly eliminated in the urine and to some extent in the fæces, it has been found in the saliva.

The **symptoms** of fatal poisoning by laburnum flowers, seeds, bark, wood, or root supervene in from five minutes up to an hour or more. They comprise—A hot feeling in the throat, thirst, vomiting, eructations, pain in the stomach, diarrhœa, collapse, cold moist surface, feeble, irregular pulse, gasping respiration, headache, drowsiness, profound prostration and coma, in some cases delirium, elevation of temperature, and convulsions resembling those due to strychnine, have occurred. The pupils are usually dilated, but they have been observed to be contracted. Death is commonly due to asphyxia, and may be preceded by cyanosis.

Poisoning by laburnum is most frequent in children who are tempted to chew or eat parts of the tree on account of its sweetish taste, out of 155 cases collected by Falk 120 were in children. An instance of wholesale poisoning resulted from 58 boys chewing pieces of the root of a laburnum tree which had been recently cut across, in the worst cases vomiting occurred with slowing of the pulse, irregular dilatation of the pupils, unconsciousness, and convulsive movements of the legs, followed by profound sleep, in all the cases the pupils were dilated, and the symptoms were of a purely narcotic type, the patients all recovered<sup>2</sup>. A case is recorded by Johnson<sup>3</sup> of six children, from eight to ten years of age, who ate laburnum seeds, they perspired and then went cold and shivery, and vomited, the pulse was scarcely perceptible at the wrist, the pupils dilated, giddiness, drowsiness, and collapse were observed. One child was purged once, and another repeatedly—in this case purging was the chief symptom, the rest were not purged, they all recovered. Two children, respectively aged three and eight years, presumably ate some laburnum seeds or pods. Vomiting, diarrhœa, and prostration occurred in one, with death in fourteen hours, the younger child felt tired and sleepy, and then vomited.

<sup>1</sup> *Freidrich's Blatter f ger Med*, 1868

<sup>2</sup> *Brit Med Journ*, 1875

<sup>3</sup> *Ibid*, 1891

and was convulsed until death occurred eight hours after the symptoms commenced. At the necropsy irritation of the gastro-intestinal mucous membrane was found, no fragments of the seeds were present in the stomach, but cytisine was detected in its contents<sup>1</sup>. Out of the 155 cases collected by Falk only four died, of the fatal cases, two had violent cramps and died within an hour, a third died in twelve hours, and the fourth not until the seventh day after taking the poison. Saake<sup>2</sup> records that of three children simultaneously poisoned by eating laburnum seeds, one a boy aged four years, died in about thirty hours, the other two, aged respectively three and four years, recovered. In all three the symptoms, singularly delayed, began with sudden vomiting and purging—in the fatal case not until fourteen hours, and in the other two not until twenty-four hours after the seeds were eaten.

**Treatment** consists in thoroughly washing out the stomach, or in giving emetics followed by copious draughts of warm water. Warm applications and friction to the surface, artificial respiration, strong coffee, and stimulants may be necessary.

The **post-mortem appearances** are negative. The signs of inflammation of the gastric mucous membrane which the symptoms during life would indicate have not always been found after death. The mucous membrane of the alimentary tract has been found pale, a few ecchymoses being present in the stomach, on the other hand, the brain and its membranes may be much congested.

**Chemical Analysis**—Cytisine is best extracted from an aqueous solution by chloroform. Radziwillowicz<sup>3</sup> recommends amyl alcohol for this purpose, but Moer and Plugge<sup>4</sup> state that the pure alkaloid is much more soluble in chloroform.

**Tests.**—Cytisine dissolves in concentrated sulphuric acid without undergoing change of colour, on warming, the mixture becomes yellow. If to a little cytisine dissolved in a few drops of concentrated sulphuric acid in the cold a drop of nitric acid is added a yellow colour is produced. If to a mixture of cytisine and sulphuric acid a fragment of potassium dichromate is added, a yellow colour is produced which changes to dirty-brown, and finally to green. With a solution of a ferric salt cytisine yields a red colour, which disappears on the addition of a few drops of a solution of peroxide of hydrogen, on subsequent warming a blue colour is produced, this test is very delicate, according to Moer and Plugge it will indicate the presence of 0.5 mgrm. of the alkaloid.

## MEZEREON.

**Daphne Mezereum**, or mezereon, occasionally gives rise to accidental poisoning in children who pluck and eat the berries, the juice being strongly irritant tends to destroy mucous surfaces with which it comes in contact.

The **symptoms** are illustrated by the following cases.—Eager<sup>5</sup> saw a child, four years old, after it had eaten at least twelve mezereon berries. Convulsions occurred before any other symptoms, an emetic was given, and vomiting procured, three hours after, the lips and tongue were swollen, the tongue, twice its natural size, was raw, and protruded beyond the lips, there was difficulty in swallowing, the limbs were cold and the pulse—130 in the minute—was very weak, recovery took place. Dunne<sup>6</sup> saw a child of the same age which had also eaten some mezereon berries. It was restless and complained of pain in the mouth and throat, vomiting took place spontaneously before the child was seen, an emetic was afterwards given, which brought away further portions of the berries. The child was drowsy,

<sup>1</sup> *Brit. Med. Journ.*, 1882.

<sup>2</sup> *Deutsche med. Wochenschr.*, 1895.

<sup>3</sup> *Ueber Nachw. u. Wirk. des Cytisins*, Diss., 1887.

<sup>4</sup> *Arch. der Pharm.*, 1892.

<sup>5</sup> *Brit. Med. Journ.*, 1887.

<sup>6</sup> *Ibid.*, 1890.



prostrate, pale in the face, with dilated pupils, scarcely perceptible pulse, and cold limbs, the mucous membrane of the tongue and of the roof of the mouth was white from the action of the acrid juice of the berries, recovery took place

**Treatment**—Evacuate the stomach, and afterwards administer an aperient with such further treatment as the symptoms require

## OIL OF TURPENTINE.

**Symptoms**—A poisonous dose of turpentine oil causes a burning sensation in the mouth and stomach, followed by symptoms of gastro enteritis. Vomiting, thirst, diarrhoea, tympanites, and a condition like that of the early stage of alcoholic intoxication are present, the pulse and respiration vary, the surface is cold, and, in fatal cases, coma supervenes, there may be muscular spasms. Strangury is a constant symptom, and the urine has an odour resembling that of violets, a similar odour being often observable in the breath, severe pain in the loins, with hæmaturia, may be present. Turpentine is excreted by the lungs, kidneys, and skin. The urine excreted after poisonous doses of turpentine has been found to reduce Fehling's solution.

Prolonged inhalation of the vapour of turpentine produces toxic symptoms which are occasionally observed in those who have slept in newly painted rooms. A case, illustrative of poisoning by the vapour of turpentine, is recorded by Reinhard<sup>1</sup>. A man who was occupied in a room in filling small vessels out of a large vessel containing turpentine, began to feel dizzy on the first day, on the second day dryness of the mouth and depression came on, and on the third day increased heaviness and painful micturition. When seen, the patient was very drowsy, the bladder was distended to the umbilicus, and the urine contained blood and albumin, it had an odour of violets which it continued to yield for a week after the patient ceased to inhale the turpentine vapour.

**Fatal Dose**—A tablespoonful has caused the death of an infant five months old, whilst another infant recovered from four ounces. To adults six ounces were fatal in one case, and in another four ounces caused death in twelve hours. Recovery in an adult has followed an ounce and a half. A case is recorded by Cardile<sup>2</sup> on which recovery followed four fluid ounces, inflammation of the intestinal tract, salivation, drowsiness and nephritis occurred. The urine contained albumin and bile pigment, but no reducing substance.

**Treatment**—The stomach pump, or an emetic, will be required, followed by demulcents. A purge should be given if diarrhoea has not occurred. Opium and other general treatment may be advisable.

**Post-mortem Appearances**—The blood has been observed to be dark coloured, and hæmorrhagic spots have been found in the stomach, sometimes with erosion of the mucous membrane.

## OIL OF EUCALYPTUS.

Oil of eucalyptus, the oil distilled from the leaves of *Eucalyptus globulus*, the blue gum tree, contains several isomeric terpenes of the formula  $C_{10}H_{16}$  possessing toxic properties. Kirkners<sup>3</sup> saw a man of twenty-eight who, after sucking numerous eucalyptus lozenges for a cold, took by mistake two or three teaspoonfuls of the oil. In a few minutes he was seized with giddiness, faintness, and ataxic gait. Acute dyspnoea, weak thready pulse, and subnormal temperature ( $95^{\circ}$  to  $96^{\circ}$  F) followed. Violent vomiting occurred, the skin became of a greenish-yellow hue, and he complained of a girdle like constriction round the abdomen. Half an hour after taking the oil he had violent diarrhoea with excessive and painful micturition. He became drowsy, and complained of intense headache, the mental faculties were blurred. Under treatment he improved, but drowsiness continued for three days, and the urine, faeces, and skin smelt strongly of the oil for nearly a fortnight. The same observer describes a case presenting similar symptoms in a girl of eighteen who had taken about a drachm of eucalyptus oil for toothache. Allan<sup>4</sup> saw a child of one year and eight months who had been given a teaspoonful of the oil by the mouth. She vomited violently and became collapsed and semi-unconscious. The temperature was  $96.4^{\circ}$  F. Complete recovery occurred in two days.

<sup>1</sup> *Deutsche med. Wochenschr.*, 1887.

<sup>4</sup> *Chin. Med. Ital.*, 1899.

<sup>2</sup> *Brit. Med. Journ.*, 1910.

<sup>3</sup> *Ibid.*

### SAVIN.

**Juniperus Sabina**, or savin, is a coniferous plant containing, as a toxic principle, an essential oil possessing a peculiar and easily recognisable odour like that which is given off by the plant, both the leaves and the oil have an acrid burning taste. Savin is rarely taken for the purpose of committing suicide, but, being regarded by the lower classes as an emetic, death has frequently resulted after it has been taken in order to procure abortion. Savin possesses no emetic properties, it is an irritant, and when abortion has ensued after its administration, the result has been due to general disturbance of the system, and not to any specific action of the poison on the womb.

The **symptoms** comprise a burning sensation from the throat to the stomach, colicky pains in the abdomen, vomiting, purging, and strangury. Labouring respiration may occur, followed by unconsciousness, collapse, coma, and death, blood may be present in the motions. It is very exceptional for abortion to take place without the woman paying the penalty with her life, on the other hand, death of the woman has frequently occurred without abortion being produced.

The **post-mortem appearances** are limited to signs of inflammation of the mucous membrane of the stomach and bowels, with the possible presence of fragments of the leaves. Sometimes no signs of inflammation have been visible, at others, punctiform ecchymoses in the gastric mucous membrane have been observed.

### YEW.

**Taxus Baccata** or common yew, another coniferous plant, owes its toxic action to the presence of an alkaloid—**taxine**—which is most abundantly present in the leaves, and to a lesser degree, in the seeds of the fruit, it is soluble in alcohol and ether, and feebly so in water.

Poisoning occurs from the use of the leaves as an emmenagogue, or as an emetic, or from accidental causes. As a uterine stimulant yew, like savin, is inert, nevertheless the working classes make use of the leaves from time to time to determine menstruations, or to procure abortion.

**Symptoms**—Giddiness, vomiting, muscular weakness, pains in the stomach and bowels, irregular action of the heart, labouring breathing, collapse, general spasms or convulsions, and delirium have been observed. A girl, on four consecutive mornings, drank a tumblerful of a decoction of yew leaves to promote menstruation, vomiting occurred, and death, preceded by delirium, took place eight hours after the last dose, the post mortem appearances were negative.<sup>1</sup> Taylor relates the case of a lunatic woman who, whilst preparing evergreen decorations, ate a few pieces of yew leaves, she became collapsed, and died in less than three hours after the symptoms first appeared, the fragments of leaves in the vomit and in the contents of the stomach after death amounted to less than a teaspoonful.

In a case (communicated by Barling) a young woman in the third month of gestation drank a teaspoonful of a strong decoction of the leaves of the Irish yew three times a day for a fortnight. After the third day, a tingling sensation was experienced in the epigastrium, which developed into severe abdominal pain, accompanied by nausea, incessant vomiting and diarrhoea. At the end of the second week the patient became unconscious for several hours, when the decoction was stopped, and the symptoms subsided in a few days without abortion taking place. In due time, the patient was delivered of a full term, dead child.

**Treatment**—Evacuate the stomach, give stimulants, relieve the bowels, and apply external warmth, with general treatment of prominent symptoms.

**Post-mortem Appearances**—Fragments of leaves or of the seeds may be found in the stomach, with signs of inflammation of the gastric mucous membrane. In a case recorded by Carter,<sup>2</sup> a girl was found dead in bed with a history leading to the assumption that she had taken yew leaves as an abortifacient, no vomiting occurred, and death took place within nine hours, the stomach contained fragments of the leaves, and the mucous membrane was inflamed.

### PENNYROYAL.

**Hedeoma**, or pennyroyal, much used as an emmenagogue, contains a volatile oil which may produce toxic effects. Wingate<sup>3</sup> states that a pregnant woman, aged twenty, took a teaspoonful of oil of pennyroyal as an emetic, she became unconscious, the limbs were cold, the pulse was small, and the pupils were slightly dilated, vomiting, delirium, and two

<sup>1</sup> *L'Imparziale*, 1870.

<sup>2</sup> *Brit Med Journ*, 1884.

<sup>3</sup> *Boston Med and Surg Journ.*, 1889.

attacks of opisthotonos occurred. Recovery, without abortion, took place. Flynn<sup>1</sup> saw a woman one hour after she had swallowed three drachms of essence of pennyroyal with the object of procuring abortion. The patient was very excited, the pupils were extremely dilated, and the pulse was feeble and became imperceptible at the wrist, after an emetic improvement rapidly set in, followed by recovery. Allen<sup>2</sup> reports the case of a woman aged twenty three who had persistent vomiting for more than four days and died on the eighth day after taking, as she stated, one tablespoonful of oil of pennyroyal. After death the stomach and the small intestines were found intensely congested, especially at the lower part of the ileum, the large intestines down to the rectum were also congested, but in a lesser degree.

### SAFFRON.

The dried stigmata of the *Crocus Sativus*, or *Saffron*, contain a volatile oil which possesses toxic properties. Occasionally saffron is resorted to as an emetic. Corvey<sup>3</sup> records the case of a woman who swallowed an unknown quantity of saffron in order to procure abortion, she became unconscious, with dilated pupils, nystagmus and paralysis of the limbs, abortion occurred in forty hours and death eight hours afterwards. At the autopsy extreme hyperæmia of the mucous membrane of the stomach with numerous submucous hæmorrhages, there, and in the intestines, were discovered, indications of slight nephritis were also present.

### TANSY.

*Tanacetum Vulgare* or tansy, contains a volatile oil, which, along with the leaves of the herb itself, has a reputation as an emetic, emmenagogue, and also as an anthelmintic. It is poisonous, and has caused death after being taken for the above named purposes. In a case recorded by Jewett<sup>4</sup> the symptoms were as follows:—A woman, aged twenty-nine, took fifteen drops of oil of tansy at eleven in the forenoon, three hours afterwards she took a teaspoonful, having had dinner between the two doses. Fifteen minutes after the second dose she threw herself on the sofa and then sprang up with a wild cry and was convulsed, respiration was for a time suspended and she became deeply cyanosed, especially about the face, neck, and hands, the eyes were open, the pupils were widely dilated, and there was great restlessness. The surface was cold and moist, the pulse was 120, and the respirations 35 to the minute, the odour of the oil being perceptible in the breath and also in the matter which the patient vomited after taking an emetic, recovery took place. In a case communicated by Dalton a woman was found on the floor in violent convulsion, she was unconscious, the cheeks were flushed and of a bright red colour, the eyes were open and very brilliant, with widely dilated and fixed pupils, the respirations were hurried, laboured, and stertorous, the breath having the odour of tansy, the pulse (128) was full. Spasms occurred at intervals by which the head was thrown back, the arms were raised and rigid, and the fingers spastically contracted, the pulse gradually grew feeble and suddenly ceased three quarters of an hour after the first appearance of the symptoms. At the necropsy no indications were discovered except the odour of tansy oil, which pervaded the entire body, and was perceived as the cavities were respectively opened, globules of the oil were found in the stomach, a fœtus at about the fourth month was found in the uterus. There was reason to believe that about eleven drachms of oil of tansy were taken.

### OIL OF WINTERGREEN.

*Oil of Wintergreen*, or oil of gaultheria, consists for the most part of methyl salicylate, it has an agreeable odour and a sweetish taste.

The symptoms produced by poisonous doses may be gathered from the following cases—Hamilton<sup>5</sup> saw a woman after she had swallowed half an ounce of the oil, she was dizzy, drowsy, and delirious. An emetic caused evacuation of the contents of the stomach, which were coated with a film of the oil and contained shreds of mucous membrane. The pupils were contracted, the respirations quick and laboured, and the limbs cold, hallucinations of audition and vision, pain in the head, and a strong disposition to sleep, verging towards coma, were present. Hemiparesis of the left side, with extreme irritability of the nervous system—starting at the least sound—and profuse salivation were prominent symptoms,

<sup>1</sup> *Brit. Med. Journ.*, 1893.

<sup>2</sup> *The Lancet*, 1897.

<sup>3</sup> *Dissert.*, Leipzig, 1895.

<sup>4</sup> *Boston Med. and Surg. Journ.*, 1880.

<sup>5</sup> *New York Med. Journ.*, 1875.

recovery slowly took place Pinkham<sup>1</sup> reports the case of a woman who swallowed one ounce of oil of gaultheria, which caused profuse sweating, pain in the abdomen, frequent painful micturition and purging, followed by convulsions, loss of sight and hearing, flushed face, rapid respirations, feeble pulse, and death in fifteen hours At the necropsy the blood was found black and fluid, and the mucous membrane of the stomach and duodenum intensely congested, the contents of the stomach yielded the odour of the oil Three drachms of the oil killed a boy three years of age, the symptoms included opisthotonos, clonic spasms, quick pulse, slow respiration, pain in the stomach, and retraction of the head

### JABORANDI.

**Jaborandi**, the dried leaflets of the *Pilocarpus pennatifolius*, owes its toxic properties to the presence of the alkaloid pilocarpine, with possibly two others

**Pilocarpine** ( $C_{11}H_{16}N_2O_2$ ) is a colourless, syrupy liquid, without odour, with acids it forms crystallisable salts, of which the hydrochloride and the nitrate are most used Pilocarpine powerfully stimulates almost all the secretions, especially the saliva and the sweat, it also stimulates the motor nerves of the involuntary muscles In small doses it stimulates, and in large doses it paralyses the vagus-endings in the heart, and thus slows or accelerates the pulse Respiration is impeded by excessive secretion of mucus into the bronchi, the temperature is lowered, the pupils are contracted and increased peristalsis of the intestines occurs Pilocarpine is eliminated in the urine

**Symptoms.**—The following case by Fuhmann<sup>2</sup> illustrates the effects exceptionally produced by a medicinal dose A man, aged thirty-one, had 01 grm (about one-sixth of a grain) of pilocarpine injected subcutaneously, the face immediately grew red, then the neck, and shortly after the whole body, which began to perspire freely In a few minutes the patient experienced sudden severe oppression of the heart and extreme difficulty of breathing, as though the chest was filled with fluid, the severe cardiac oppression subsided in ten minutes, but traces could be felt for two hours Increased secretion of saliva, tears, and of mucus from the nostrils occurred Cramps were experienced in the stomach as though the organ "would turn round", nausea and vomiting followed, with movements of the intestines which produced a strong desire to evacuate the bowels Great thirst, prostration, and a tired feeling, especially in the legs, were experienced, the pupils were contracted and vision was impaired, the pulse was small and frequent, and the patient became collapsed The amblyopia lasted two hours, the sweating two and a half hours, the salivation four and a half hours, during which time 500 grammes of saliva were expectorated, the patient recovered A case is recorded by Sziklai<sup>3</sup> in which a patient by mistake had fifteen minims of a 20 per cent solution of pilocarpine injected under the skin Immediately on withdrawing the cannula, salivation, copious sweating, and shortly after dizziness, vomiting, diarrhoea, oppression, and a tearing pain in the eyeballs with pronounced myopia occurred, the pupils were contracted to the uttermost The patient micturated twice, the salivation and diaphoresis lasted two hours, after five hours the acute symptoms began to disappear, and the patient recovered The subcutaneous injection of from a thirtieth to a sixth of a grain of pilocarpine hydrochloride caused the outbreak of a papulous exanthema followed by death (Hallepeau and Viellard)<sup>4</sup> Remy<sup>5</sup> relates a case in which, after a subcutaneous injection of pilocarpine—dose not stated—the patient fell back dead Jaborandi has also produced psychical disorder with hallucinations

<sup>1</sup> *Boston Med. and Surg Journ.*, 1888

<sup>2</sup> *Wiener med Wochenschr.*, 1890

<sup>3</sup> *Wiener med. Wochenschr.*, 1881.

<sup>4</sup> *Ann de Dermat et Syphilogr.*, 1905.

<sup>5</sup> *Recueil d'Ophthal.*, 1893.

**Treatment.**—If necessary evacuate the stomach and then give stimulants. One-fiftieth of a grain of atropine sulphate should be injected subcutaneously, and repeated if necessary.

### CALABAR BEAN.

**Calabar Bean**, the seed of the *Physostigma venenosum*, is kidney-shaped, having a groove with elevated borders along its convexity, it is blackish-brown in colour, and its surface has a texture like that of the fine morocco leather used for bookbinding. The bean varies from an inch to an inch and a half in length, if split lengthwise it is seen to consist of a brown-coloured rind containing two white cotyledons which adhere to the shell. It contains two alkaloids—physostigmine and calabarine, its characteristic toxic effects being due to the former.

**Physostigmine** ( $C_{15}H_{21}N_3O_2$ ), or **eserine**, when pure, is a white crystalline body which rapidly changes colour on exposure to air and light, it is not very soluble in water, but is easily soluble in alcohol, ether, and chloroform. It has an alkaline reaction, and forms salts with acids, which are colourless and soluble in water, on exposure to air they deliquesce and acquire a red colour.

Physostigmine increases the irritability of both the voluntary and the involuntary muscles—shown by twitchings of the skeletal muscles and peristalsis of those of the intestines. It eventually paralyses the respiratory centres in the medulla and causes death from arrest of respiration. It augments the irritability of the vagus, and thus tends to slow the heart, it increases the secretions, probably by direct action on the secreting elements themselves, and paralyses the motor centres in the brain and cord. When applied directly to the eye, physostigmine contracts the pupils and causes spasm of accommodation, probably from stimulation of the third nerve. Physostigmine is eliminated in the urine, faeces, and saliva.

**Calabarine** resembles strychnine in its action, and causes clonic spasms.

**Symptoms.**—After a poisonous dose of calabar bean, giddiness and faintness are experienced, quickly followed by profound prostration, pain is felt in the stomach, usually followed by vomiting, diarrhoea may occur, but is not very frequent. The heart's action is enfeebled, the pulse being usually small and slow, and there may be dyspnoea. The surface is cold and moist, the pupils may be, but are by no means invariably contracted, salivation and thirst have occurred. The mental condition varies in some instances it is undisturbed, in others drowsiness and even unconsciousness have been observed. Muscular spasms have been present in rare cases, probably due to calabarine. In the year 1864, forty-six children were poisoned in Liverpool through eating calabar beans which had been discharged from a ship, they all suffered from pain in the region of the stomach, thirty-eight were attacked with vomiting, and fifteen with diarrhoea, only one died.

A unique case of suicidal poisoning with physostigmine is recorded by Leibholz<sup>1</sup>. Two girls, aged twenty-four and eighteen respectively, obtained possession of a sealed tube containing 0.1 grm of physostigmine sulphate, which they dissolved in water, and each girl drank half of the solution, for half an hour they pursued their household avocations without experiencing any effects, they then suddenly became unconscious. In each case the face was red and shining, the pupils, dilated to the maximum, were reactionless,

<sup>1</sup> *Vierteiljahrsschr. f. ger. Med.*, 1892.

the pulse, 60 to the minute, was full and of high tension, the respirations were shallow rapid, and moaning, pain was experienced in the region of the stomach and abdomen, vomiting occurred early, and persisted for some time after return to consciousness. Dilatation of the pupils, with feeble reaction to light, lasted for several days, perfect recovery ultimately taking place. The activity of the alkaloid was vouched for by Merck after chemically examining a companion sample. Tested physiologically three milligrammes injected under the skin of a rabbit weighing four pounds produced paralysis of the voluntary muscles, difficulty of respiration, violent diarrhœa, and death in ten minutes, a solution dropped into the human eye caused marked contraction of the pupil. The dilatation of the pupils in the above recorded cases is remarkable, instances of calabar bean poisoning have occurred without contraction of the pupils, but, with the above exception, none with dilatation. The absence of diarrhœa is in marked contrast to its universal occurrence in animals poisoned with physostigmine.

Eserine (physostigmine) has produced symptoms of poisoning after being dropped into the eyes for ophthalmic purposes. Dunlop<sup>1</sup> relates the case of a man, aged sixty, who had one drop of a solution of eserine (inadvertently prepared with one grain to the drachm instead of to the ounce) dropped into each eye, a quarter of an hour after clonic spasms of the eyelids, stiffness of the lips, a feeling of tremor in the arms and legs, together with mental confusion occurred, the symptoms passing off after a time. A one per cent solution of physostigmine saleylate sprayed into the nose caused facial pallor, weak action of the heart, cold perspiration, and great anxiety (Speer).<sup>2</sup>

**Treatment.**—If the poison has been swallowed evacuate the stomach with the tube or an emetic, then give stimulants, and apply external warmth. Atropine sulphate injected subcutaneously in doses of one-fiftieth of a grain, repeated until the pupils dilate and the pulse-rate is quickened, is recommended by Fraser, strychnine is also regarded as an antidote. Should respiration fail, it must be promoted artificially.

**Post-mortem Appearances** are negative

**Chemical Analysis.**—The alkaloid may be extracted from organic matter in the usual way, special care being taken not to allow excess of acid, heat, or light to come into play, ether is a good solvent.

**Tests.**—Bromine water, added to a solution of physostigmine, produces a red or orange-coloured turbidity, which clears up on heating. Strong chlorine water added to the solid alkaloid produces a red colour. When physostigmine is treated with a little ammonia and the mixture is evaporated to dryness, the residue is of a more or less intense blue colour, often with a reddish appearance in parts. When very little of the alkaloid is present, the colour will be greenish. If the blue deposit is dissolved in alcohol and is examined with the spectroscope it yields a band in the red, on acidulating the solution the blue changes to red and gives an absorption band in the yellow. If a solution of barium oxide in water is added to urine containing physostigmine, or to an aqueous solution of the alkaloid, a white precipitate forms which on boiling and shaking becomes red.

The physiological test should be resorted to — a drop of an aqueous solution instilled into the eye of a rabbit produces contraction of the pupil, the test may be repeated on the human eye.

<sup>1</sup> *The Lancet*, 1887.

<sup>2</sup> *Therap. Gaz.*, 1905.

## NUTMEG.

**Nutmeg**, the kernel of the seed of the *Myristica fragans*, though of every day use as a food adjunct, is capable of causing poisonous symptoms when taken in excess. Bentliff<sup>1</sup> records the case of a man who swallowed the whole of a grated nutmeg as a remedy for boils. He afterwards went to bed and slept until six o'clock in the morning, when he felt giddy and could not stand, he had pain in the head, and could not distinguish objects, he was drowsy, but could be roused by shouting, thirst, numbness of the limbs, and some what contracted pupils were the other symptoms present, he recovered in a short time. Sawyer<sup>2</sup> relates the case of a boy, aged three, who ate portions of five nutmegs. He was dizzy, and then became unconscious, with relaxed muscles, and could not be roused, he slept for thirty consecutive hours, the pupils were dilated, there was no delirium. In contrast to this case is one related by Reading,<sup>3</sup> in which a lady, three months pregnant, in order to procure abortion, swallowed three powdered nutmegs, she vomited several times, and became delirious with hallucinations, accompanied by laughter, the pulse was strong and rapid. Notwithstanding treatment, the delirium recurred at intervals during the succeeding twenty four hours, when she became rational, and recovered without abortion taking place.

## CAMPHOR.

**Camphor** in poisonous doses acts as an irritant and produces the usual symptoms of collapse, it first stimulates and then paralyzes the nerve centres, and tends to cause convulsions. East<sup>4</sup> records the case of a youth, aged nineteen, who swallowed a large lump of sugar saturated with essence of camphor, and an hour afterwards drank an equal quantity added to some water, the total amount being about two drachms. In two hours after the first dose he felt giddy, talked incoherently, and fell down in a "fit". The pulse was full, and the pupils were dilated, the patient vomited spontaneously, the rejected matter having the odour of camphor, recovery took place. Davis<sup>5</sup> saw a child, two and a half years old, after it had eaten a piece of solid camphor the size of a nut—about half a drachm, it was pale and convulsed, the lips being blue, and the pulse very rapid, on evacuation of the stomach it recovered somewhat from the collapse, but died in eighteen hours after swallowing the poison. Honman<sup>6</sup> saw a girl of eighteen who had swallowed an unknown dose of camphor, she was unconscious, the pupils were dilated, the legs were cold, the pulse was thready, and the breath had the odour of camphor, convulsive movements occurred, followed by paralysis, the breathing became shallower, and the patient died within thirty-six hours, at the necropsy camphor was found in the stomach. A child, five years of age, died in convulsions shortly after swallowing a teaspoonful of camphorated oil, whilst another child of exactly the same age recovered after swallowing a teaspoonful of crude camphor oil containing between one and two hundred grains of camphor which produced convulsions and collapse, the child vomited freely under the influence of an emetic and the next day was quite well (Wilkinson).<sup>7</sup>

## SANTONIN.

**Santonica**, the dried flower heads of the *Artemisia maritima*, contains a crystalline active principle **santonin**, which is almost insoluble in water, but is freely soluble in hot alcohol, and in chloroform. Cases of poisoning by santonin, or by substances containing it, occur almost exclusively among children, to whom it is given as a vermicide.

**Symptoms**—The most constant symptom is one that occurs even with non-poisonous doses—disturbance of colour-vision. Objects first assume a bluish tinge, and subsequently yellowish green or yellow. The most probable explanation is that primary stimulation of the violet seeing retinal elements takes place—causing violet or blue vision, with sequential paralysis and consequent absence of blue vision, the period of stimulation being very short may pass unobserved, the paralysis lasts much longer, and, therefore, the condition is usually described as "yellow vision". Singing sounds in the ears, dizziness, pain in the stomach, vomiting, convulsions, and stupor, with a tendency to asphyxia, are among the remaining symptoms, the urine is coloured saffron yellow. In a severe case observed by Demme<sup>8</sup> a boy, aged three, took about two thirds of a grain of santonin six times a day

<sup>1</sup> *Brit. Med. Journ.*, 1889.<sup>2</sup> *New York Med. Journ.*, 1889.<sup>3</sup> *Therap. Gaz.*, 1892.<sup>4</sup> *Brit. Med. Journ.*, 1886.<sup>5</sup> *Ibid.*, 1887.<sup>6</sup> *Australian Med. Journ.*, 1888.<sup>7</sup> *Brit. Med. Journ.*, 1898.<sup>8</sup> *Klinische Mittheilungen*, 1891.

On the third day there was vomiting, dilatation of the pupils, coldness of the surface, cyanosis of the lips and cheeks, dyspnoea and convulsions, with deep stupor. Bleeding at the nose and hæmoglobinuria occurred, the urine had a dark saffron tint with a greenish shimmer, the temperature was 102.5° F., cold effusion was used, and recovery took place, a scarlatina-like rash appearing on the chest.

**Chronic Santonin poisoning** is very unusual. Roy<sup>1</sup> records the case of a boy, aged eleven years, who, on account of pain in the abdomen, which was supposed by his mother to be due to worms, had santonin given to him for months. Clonic spasms then developed, to combat which the doses of santonin were increased. Paralysis, twitchings, dizziness, pain in the head, vomiting, yellow and violet vision, sparks before the eyes, and finally loss of speech occurred, necessitating medical advice, under treatment the patient was able to walk in six weeks, but it was nine weeks before he regained the power of speech.

**Fatal Dose**—About two grains of santonin taken twice proved fatal in twelve hours to a child five and a half years old, recovery in a child has followed ten grains. A man aged forty took one ounce in mistake for Epsom salt, giddiness, incessant vomiting, prostration, laboured breathing, and epileptiform convulsions ensued, but recovery eventually took place.<sup>2</sup>

**Treatment**—Evacuate the poison and administer stimulants. Convulsions may be combated by potassium bromide and chloral hydrate.

**Post-mortem appearances** are not characteristic. In a case recorded by Kilner<sup>3</sup> the stomach and duodenum of a child aged four and a half, who died thirty five minutes after taking six grains of santonin, displayed signs of inflammation.

**Chemical Analysis**—Santonin may be extracted from acid aqueous solutions by shaking out with chloroform, it will not come away from alkaline solutions, as it plays the part of a weak acid, forming combinations with alkalies which are soluble in water.

**Tests**—A solution of sodium hydroxide produces a violet red colour with santonin. Dragendorff has devised a modification of a former test which is thus performed.—A little sulphuric acid diluted with half its volume of water is added to some santonin, and gently heated until a yellow colour is produced, when cold a few drops of a very dilute solution of ferric chloride are added, and on again warming a blue or reddish violet colour is produced.

The presence of santonin in the urine may usually be ascertained by the addition of a little sodium hydroxide, if present, a red colour is produced. Rhubarb present in the urine yields the same reaction, but if, after the addition of sodium hydroxide, excess of milk of lime is added, the urine afterwards being filtered, the filtrate is colourless if the reddening is due to rhubarb, but retains its colour if it is due to santonin.

## COLOCYNTH.

**Colocynth**, or bitter apple, the dried pulp of the *Citrullus colocynthis*, contains an active principle **colocynthin**, which is soluble in water and alcohol but not in ether. Colocynthin is an active cathartic, and, in large doses, a gastro intestinal irritant. A few fatal cases of poisoning by colocynth are recorded, a teaspoonful and a half proved fatal in twenty-four hours. Tidy<sup>4</sup> states that a woman took from one to two drachms of bitter apple as an emmenagogue, on the following day, vomiting and violent purging occurred, and the patient died in about forty hours after taking the poison, the result of the post mortem examination was negative. In one case the intestines were found reddened, and the stomach ulcerated, the kidneys and bladder being inflamed.

## ELATERIUM.

**Elaterium**, the dried juice of the *Ecballium elaterium*, contains an active principle **elaterin**, which is insoluble in water but is soluble in hot alcohol and in chloroform. Elaterium is an exceedingly powerful cathartic, and in poisonous doses may produce, in addition to its drastic effects, nausea, eructation, vomiting, salivation, prostration, clonic spasms, insensibility, and dyspnoea. Two fifths of a grain has proved fatal. One sixth of a grain caused the death of a woman aged seventy. Recovery has taken place after three quarters of a grain of elaterin, severe symptoms of poisoning being produced.

<sup>1</sup> *Therap. Monatshefte*, 1889.

<sup>2</sup> *Annali univ. di Med.*, 1882.

<sup>3</sup> *St. Thomas's Hosp. Reps.*, 1880.

<sup>4</sup> *The Lancet*, 1868.



## CROTON OIL.

**Croton Oil** is a fixed oil expressed from the seeds of the *Croton tiglium* which resemble, but are smaller than, castor oil seeds. Both seeds and oil are active gastro intestinal irritants, by direct contact, the oil produces inflammation of the skin, as well as of mucous membranes.

**Symptoms**—When taken in poisonous doses into the stomach, croton oil produces a hot, burning sensation in the mouth and throat, pain in the stomach and abdomen, violent vomiting and purging, with dizziness, great prostration, cold surface and collapse. The pulse and the respirations are slowed, in fatal cases, death takes place in a few hours. Half a drachm of the oil in one instance, and twenty drops in another, proved fatal. Recovery has taken place after a drachm, and even after half an ounce, but in the latter case the oil was not pure. The seeds have also proved fatal, in one case four caused death.

## CASTOR OIL SEEDS.

**Castor Oil seeds**, the seeds of the *Ricinus communis*, are smooth, oval shaped and bean-like, in addition to the oil, which is a well known harmless purgative, the seeds contain a poisonous phytalbumose—Ricin—which Stillmark<sup>1</sup> regards as an unformed ferment. From experiments on animals it is found to be a gastro intestinal irritant, it possesses no intrinsic purgative properties, hence in some case of poisoning by castor oil seeds, purging has been conspicuous by its absence.

**Symptoms**—They comprise nausea, pain in the stomach, with a burning sensation in the throat, persistent violent vomiting, colicky pains in the abdomen, pale, collapsed appearance of the face, cold surface, great prostration, small pulse, with retention of consciousness in some cases and insensibility in others. Purging may or may not occur.

In a case recorded by Langerfeldt,<sup>2</sup> a boy ate ten to fifteen seeds, he vomited—the rejected substance containing blood—had headache, and lay on his back groaning, with his legs drawn up, pale, cold, and cyanosed, the skin was clammy, the pulse (110) scarcely to be felt the abdomen was retracted, and the tongue was dry and furred. He complained of a burning sensation in the throat and pain in the epigastrium, the bowels were obstinately constipated, on the sixth day he was quite well. F. J. Smith<sup>3</sup> records the case of a man aged twenty five who, after eating about twelve castor oil seeds, had violent abdominal pains and purging, with vomiting and frequent micturition, he was collapsed and had cramps in the calves. When the abdomen was stroked the muscles went into violent tonic contractions, recovery took place. In a case recorded by Bouchardat,<sup>4</sup> a girl, aged eighteen, ate about twenty seeds, which produced violent purging, vomiting, and profound collapse, the stools consisted chiefly of blood stained serous fluid, death occurred on the fifth day. The mucous membrane of the stomach was found softened and abraded in parts. Meldrum<sup>5</sup> records the case of a man, aged twenty six, who died six days after eating two seeds, vomiting and purging were incessant.

**Fatal Dose**—Three seeds proved fatal in forty six hours to an adult, aged thirty two. Recovery, even in children, has followed larger doses, adults have recovered from seventeen and twenty seeds respectively. Park<sup>6</sup> records the recovery of a man who had eaten twenty-four seeds.

**Treatment in Poisoning by Cathartics**—Promote evacuation of the poison, and then administer morphine hypodermically, followed by stimulants and external warmth. As soon as the stomach will retain anything, demulcents with a little ice may be given. If purging is excessively violent, starch and opium enemata may be advisable, mustard-leaves to the abdomen and epigastrium will be of service. If the collapse is very severe, hypodermic injections of ether may be needed.

**Post-mortem appearances** are usually limited to signs of inflammation of the gastro-intestinal mucous membrane, such as hyperæmia and softening, with possibly erosion in parts. Fragments of the seeds should be sought for.

## ERGOT.

**Ergot** is a parasitic formation consisting of the mycelium of the *Claviceps purpurea* developed from the ovaries of various graminæ, especially rye, it occurs in wet seasons, and may be so widely diffused as to give rise to epidemics of ergotism in the districts where the diseased grain is grown.

<sup>1</sup> *Dissert.*, 1888.

<sup>2</sup> *Berliner klin. Wochenschr.*, 1882.

<sup>3</sup> Taylor and F. J. Smith, *Medical Jurisprudence*

<sup>4</sup> *Annales de Therapeutique*, 1872

<sup>5</sup> *Brit. Med. Journ.*, 1900

<sup>6</sup> *Glasgow Med. Journ.*, 1880.

Ergot contains more than one active principle, Kobert has found three — *ergotanic acid*, *sphaellinic acid*, and *conutine*, the last is regarded as an alkaloid. the substance known as **ergotin** is an admixture of these principles. Although by means of experiments on animals, considerable information has been obtained as to their respective actions, the specific effects produced by them on the human subject have not yet been satisfactorily differentiated, from the toxicological standpoint therefore ergot and ergotin are to be regarded as complex bodies which possess certain definite toxic properties.

Ergot poisoning may be **acute** or **chronic**, the latter is frequently named ergotism.

**Symptoms of Acute Ergot Poisoning.**—When one or more poisonous doses of ergot, or of ergotin, are taken, giddiness, pain in the stomach, thirst, nausea, vomiting, great oppression in the cardiac region, numbness and tingling—beginning in the fingers and toes, and tending to spread along the limbs—cramp, dyspnoea, shivering, coldness, especially of the limbs, great anxiety, delirium, coma, and convulsions are among the symptoms which may be produced.

In a case recorded by Debierre,<sup>1</sup> a woman recovered after swallowing one drachm and a half of Bonjean's ergotin. In a few hours intense dyspnoea, faintness, dryness of the mouth and throat, giddiness, noises in the ears, dimness of vision, tingling, and a sensation of coldness in the limbs were experienced. There was complete anæsthesia of the tongue and of the surface of the body with severe pain in the epigastrium and the abdomen, the temperature was 96.8° F, the pulse 50 per minute, and the respirations also 50 per minute, epileptiform convulsions occurred. In another and fatal case recorded by Davidson,<sup>2</sup> a woman who was pregnant had been taking the liquid extract of ergot for several months, she then swallowed "two handfuls" of powdered ergot without infusing it. When seen the day after, the face and the upper part of the body were jaundiced, ecchymoses were present under the skin around the eyes, the lips and tongue were swollen and coated with dry black blood, there was intense thirst, the skin was pale, and the temperature 96° F. The pulse was peculiar—it could not be counted, but could be just perceived, and then disappeared before its character could be estimated, the heart beats wore of a rolling character—150 per minute, the respirations were 47 per minute. The patient had periods of stupor and apathy, she vomited red pulaceous matter and pure blood, the urine also contained blood. An attempt was made to effect instrumental delivery, but the woman died before it could be accomplished, the respirations increased to 56, and stupor with paroxysmal movements supervened immediately before death. At the necropsy much fluid blood, effused from small vessels, was found extravasated in the abdominal cavity, but no large vessel was ruptured, the liver, kidneys, and lungs were bloodless, the liver and kidneys presenting a pale yellow, waxy appearance, all the viscera, though bloodless, were ecchymosed, and ruptured vessels were found within the stomach and bowels. In the uterus, which contained no blood, a five months' foetus was found. The bladder was empty.

Occasionally ergot produces toxic effects when administered medicinally. Heller\* relates two cases, in one of which five seven grain doses of powdered ergot, and in the other three fifteen grain doses daily for five consecutive days caused paræsthesiæ and anæsthesia of the right arm, in the second case there was also spasm of the fingers of the left hand with painful contraction of the muscles of the legs and thorax. Venous thrombosis has followed the hypodermic injection of dialysed ergotine.

The effect of ergot on the quiescent uterus is discussed in the section on criminal abortion.

**Treatment.**—Evacuate the stomach with the tube or an emetic, and clear out the bowels. Stimulants and external warmth will be needed. Inhalations of amyl nitrite, or, as recommended by Murrell, nitroglycerine, administered by the mouth, may be tried.

<sup>1</sup> *Bullet Gen de Therap.*, 1884.

<sup>2</sup> *The Lancet*, 1882.

\* *Dissert.*, Erlangen, 1896.

**Post-mortem appearances** chiefly consist in the presence of ecchymoses and extravasation of blood on and into the internal organs, as described in the above-quoted case. The bodies of three pregnant women who died in consequence of taking ergot to procure abortion, all presented unusual post-mortem appearances—externally they were jaundiced to some extent, internally the usual ecchymoses were found, and in addition, the liver in all three, and the kidneys in two, showed fatty changes, so pronounced as to give rise to suspicion of phosphorus poisoning. On chemical examination ergot was found in the intestines, but no trace of phosphorus. In two of these cases the uterus contained a four months' and a six weeks' fœtus respectively, in the third, a fœtus surrounded by its membranes was at the vaginal outlet.<sup>1</sup>

**Symptoms of Chronic Ergot Poisoning.**—Chronic ergot poisoning mostly occurs as a consequence of eating bread made from grain contaminated with the fungus, it has occurred epidemically for many centuries, and still appears from time to time in Germany, Russia, and other countries.

The **early symptoms** indicate disturbance of the gastro-intestinal tract, they comprise pain and oppression in the gastric region, general depression, either loss or increase of appetite, nausea, occasional vomiting, sometimes diarrhœa and at others constipation, together with dizziness, sleeplessness, and a general feeling of weariness and absence of energy. Subsequently the symptoms may take one or both of two courses—**gangrenous** or **nervous** (spasmodic ergotism).

**Gangrenous Ergotism** is first indicated by the occurrence of patches of anæsthesia—the patient experiencing a sensation of cold in the parts affected—or by a burning sensation, accompanied by redness of the skin. Gangrene, mostly of the dry type, which may or may not be preceded by the formation of serous blisters, then sets in, the peripheral parts of the limbs—the toes and fingers—being most frequently affected. The gangrene, which seldom affects the trunk, may advance as far as the knees or elbows, when it has reached its limit, separation by slow ulceration takes place, unless the process is expedited by surgical operation. In rare cases the skin only is attacked, the entire cutis undergoing necrosis, and separating from the underlying tissues.

**Spasmodic Ergotism** is preceded by paræsthesiæ of various kinds, such as a creeping sensation beginning in the fingers and toes and spreading along the limbs, in some instances there is complete anæsthesia. Motor disturbances follow—first, twitching of the muscles occurs, then spastic contraction of groups of muscles, by which the fingers and toes are flexed and drawn together, the hands are flexed at the wrists, presenting the appearance of clenched fists with the thumbs drawn towards the palms, the ankles are extended, the heels being sometimes so powerfully drawn up that the feet and the legs form straight lines. The spasm may extend along the muscles of the limbs to those of the spine so as to produce opisthotonos, rarely the muscles of the lower jaw are similarly affected. The spasms are exceedingly painful, and cause the patient to roll about in agony, the surface being covered with a cold sweat, they last from a few minutes to many hours, when they pass off the patient is left exhausted and powerless. Sometimes the contractions are tetanoid in character, at others, clonic spasms resembling epilepsy occur, the breathing may be affected as though the diaphragm participated in the spasmodic seizure. Dysuria due to spastic contraction of the bladder may be present. Paralysis, with complete superficial anæsthesia, sometimes follows. Disturbances of the special

<sup>1</sup> *Petersb med Wochenschr*, 1884

senses have been recorded, as diplopia, alterations in the field of vision for colours, deafness, and aphasia. Exceptionally cataract has been observed. Orlov<sup>1</sup> believes that the changes in the eyes are not due to spasm of the vessels, but to the immediate toxic action of the ergot on the retina and other tissues of the eye.

Psychical disorders, as hallucinations, delirium, mania, mental enfeeblement, with stupor, and, exceptionally, indications of tabes—lightning pains, girdle sensation, staggering gait and unsteadiness in the erect posture with the eyes closed—have occurred. Tuczek<sup>2</sup> found sclerosis of the posterior columns of the cord, implicating the root-zones as in tabes.

Both the gangrenous and spasmodic types of ergotism are probably due to persistent contraction of the smaller arteries, which deprives the tissues respectively implicated of their normal supply of blood, as previously stated, the two varieties may coexist—a patient with spasmodic ergotism may lose the toes and fingers from gangrene.

From a report made by Griassnoff<sup>3</sup> on seventeen cases of ergotism which were admitted into the Poltava hospital in Russia during the epidemic of 1881, the following statements are taken—The ages of the patients varied from twelve to forty-five years, thirteen were males and four were females, four died—two males and two females. All suffered from agonising pains and numbness of the limbs, sleeplessness, exhaustion, diarrhoea, weak accelerated pulse, and all but one, from loss of appetite. Five suffered from spasms, a few from headache, nausea, and vomiting. In all but one, gangrene of the humid type in eight, and of the dry in seven—occurred, all of these patients had pyrexia (101° F and more), with evening exacerbations. The quantity of ergot present in the rye meal which had been eaten by the patients was not more than one per cent, which is much lower than the percentage usually stated as liable to cause ergotism.

The treatment of ergotism is mostly prophylactic, with ordinary medical or surgical treatment of the symptoms as they arise.

**Chemical Analysis.**—Bread or flour suspected of containing ergot should be extracted with hot alcohol acidulated with sulphuric acid. The extract is red in colour, and, if examined with the spectroscope, yields two bands, one in the green and another in the blue, the latter being the broadest and best defined. It is practically impossible to separate ergot from the tissues so as to identify it, in acute poisoning, the contents of the stomach may be treated as above, and the ergot if present identified.

## LATHYRISM.

**Lathyrism**, caused by eating grain derived from some of the vetches, closely resembles the pathological condition which is developed by changes in the lateral columns of the spinal cord, tremors, spastic gait, stiffness of the muscles of the back and legs, with excessive knee-jerk and sensory disturbances. The condition subsides after the grain has ceased to be eaten.

## FUNGI.

Fungi have been classified as edible and poisonous, some are definitely known to be poisonous, but it by no means follows that all others may be eaten.

<sup>1</sup> *Neurolog Westnik*, 1905

<sup>2</sup> *Arch f Psychiat*, 1882

<sup>3</sup> *The London Med Record*, 1883

with impunity. It is obvious that those only which contain intrinsic toxic principles can be classified as poisonous, according to Husemann, these comprise—*Amanita muscaria*, *Amanita phalloides*, *Russula integra*, *Boletus luridus*, and their varieties. Such fungi are poisonous in the same sense as are any of the known poisonous vegetables, there are, however, many fungi which contain no essential toxic principle, and yet from time to time they act as poisons. In England the common mushroom (*Agaricus campestris*) and the champignon (*Agaricus oreades*) are the only fungi eaten, on the Continent a much more liberal selection is made.

The erratic way in which fungi, regarded as harmless, occasionally produce violent toxic effects, has been variously accounted for. Many edible fungi contain amanitin, which, though itself inert, may be resolved by incipient decomposition into neurin—a closely allied, or as some authorities hold, an identical substance—and may thus give rise to symptoms of poisoning. Some edible fungi contain more albumin and fatty matter than others, and are, therefore, more liable to undergo decomposition either before gathering, when past their prime, or after gathering and before cooking. The morel, according to Kohlrausch, contains 35 per cent of albumin and 2.39 per cent of fat, whereas the common mushroom contains only 17 per cent albumin and 1.4 per cent fat—hence poisoning from decomposition-changes is less frequent by the latter than by the former. It has been supposed that fungi (especially the morel) when gathered in wet weather are likely to become poisonous, in some instances it is probable that specimens of a poisonous variety have been accidentally included among edible fungi. Idiosyncrasy may have something to do with the matter, but not much, the severity of the symptoms respectively produced in a number of people who have partaken of a poisonous dish of mushrooms, depends on the quantity each has eaten, and more especially on the amount of the juice or gravy consumed (as representing an extract of the fungi) rather than on idiosyncrasy. Many years ago the author saw three fatal cases of mushroom poisoning in one family. The mother and three children partook of mushrooms for supper, and were taken ill the following morning, with symptoms of acute gastro-enteritis, the mother and two of the children died within forty-eight hours, the third just escaped. The symptoms were by far the most violent in the woman, and on inquiry it was found that after helping the children she took the dish to herself and soaked up the juice with bread, which she ate in addition to the mushrooms, the excess in number of mushrooms she ate in comparison with those eaten by the children, was sufficient alone to account for the relative violence of her symptoms. That the poison is easily dissolved out is shown by the fact that in some parts of the Continent the poisonous fly-fungus (*Amanita muscaria*) is eaten with impunity after being well extracted with water. Mushrooms which have been dried and kept for some time may develop ptomaine-like poisons, mushrooms which have been cooked, should not be eaten after being put on one side and warmed up again.

**Symptoms of Poisoning by Fungi** are divisible into two groups—*gastro-enteric* and *neurotic*—both of which are usually represented in the same patient.

**Gastro-enteric symptoms** may not appear for six or ten hours after the fungi are eaten, and not infrequently they are still further delayed. A feeling of uneasiness in the stomach gradually develops into pain, with a hard tender condition of the abdomen, nausea is experienced, and then vomiting which is followed by diarrhoea. The vomiting and diarrhoea are not solely due to the immediate presence of the irritant, but to the condition set up by it in the gastro-intestinal mucous membrane, therefore, they do not at once subside when

all the fragments of fungi are discharged, the enteric derangement is further shown by the character of the evacuations, which are serous—rice-water-like

and contain flakes of lymph, and sometimes blood, notwithstanding treatment the diarrhœa and vomiting may persist for several days. Great thirst, prostration, shrinking of the tissues, livid countenance, cold surface, small pulse, and laboured respiration are the natural results of the excessive drain on the blood, exceptionally jaundice may occur. These symptoms may directly lead to death with or without the appearance of any special nerve complications, or they may subside, and recovery may take place.

**Neurotic symptoms** comprise muscular twitchings, general convulsions or tetanic spasms, delirium, disorders of the special senses especially of vision, with dilatation of the pupils, and stupor, or profound coma. In some instances the symptoms are solely neurotic, such cases present all the appearance of certain forms of alkaloidal poisoning.

Illustrative of the gastro-enteric symptoms is the following case reported by Boyce<sup>1</sup>—A man, aged fifty-three, ate heartily of a dinner of which mushrooms formed a part, three and a half hours after he felt griping pain in the abdomen followed by diarrhœa, and on the following morning by vomiting, constant pain and vomiting continued for two days, when he sought medical assistance. His face was then dusky and bluish, the pupils were dilated, the breathing was short and rapid, the pulse weak and quick, and the surface cold, much pain was experienced in the stomach, and there was great prostration. In spite of treatment the vomiting and diarrhœa continued, the motions resembling dirty water with flocculi of lymph, death took place on the fourth day after the mushrooms were eaten. The deceased's son, who also partook of the mushrooms, was attacked in the same way, but he recovered. The following case reported by Matthes<sup>2</sup> illustrates the neurotic type of mushroom poisoning. A woman and three children were taken ill, about four hours after eating mushrooms, with pain in the abdomen and delirium. In each case the face was cold and pale, the pulse slow, the lips were cyanosed, the respirations quick and shallow, and the pupils dilated and inactive, for two hours violent clonic spasms, like those due to strychnine, occurred every eight or ten minutes, together with coma, all the patients recovered. Cases have been recorded in which the symptoms have been of a purely narcotic type.

Muscarine the active principle of the fungus *Amanita muscaria* or fly-fungus, has been isolated, and its properties have been investigated by Schmiedeberg and Koppe (who were the first to obtain it in a pure form) and subsequently by others. An infusion of the fresh fungus acts as a fly-exteriorinator—hence its name, the effect on flies, however, is not due to muscarine, which is innocuous to them, but to some other and probably volatile substance, since the dried fungus no longer acts as a fly-poison. The fly-fungus has been used by the poor in Siberia and Kamtschatka as an intoxicating medium, its active principles are eliminated by the kidneys, and this is so well known that those habituated to its use drink their own urine, or that of others who have partaken of the fungus, in order to produce intoxication.

**Muscarine** ( $C_5H_{15}NO_3$ ) is a colourless, syrupy liquid, without taste or smell. It is alkaline in reaction, is soluble in water and alcohol, slightly so in chloroform, and is insoluble in ether, it forms salts with acids, of which the nitrate is most commonly met with.

According to Brunton, muscarine produces uneasiness in the stomach, vomiting, purging, a feeling of constriction in the neck, want of breath, giddiness,

<sup>1</sup> *Brit. Med. Journ.*, 1887.

<sup>2</sup> *Berliner klin. Wochenschr.*, 1888.

faintness, prostration, and stupor. It slows the heart by stimulating the intra-cardiac inhibitory apparatus, lowers the blood pressure, depresses the respiratory centres, causes contraction of the pupils, and of the muscular coat of the intestines, stimulates the secretion of sweat and saliva, and diminishes that of urine. Muscarine strongly resembles pilocarpine in its action, and is antagonistic to atropine. Muscarine and pilocarpine differ, however, in their action on the pupils, when applied locally pilocarpine contracts the pupils, muscarine dilates them, both contract the pupils when administered internally.

So much for the physiological action of muscarine. In the human subject poisoning by fly-fungus presents other symptoms. There may be delirium, clonic spasm or convulsions, often dilated pupils, and quick pulse, this difference has led to the assumption that another active principle is present in the fungus, which is more or less antagonistic to muscarine. Harmsen<sup>1</sup> states that, in addition to muscarine, the fresh fly-fungus contains another poison which acts centrally—a fungus toxin, he holds that the toxic action of the fly-fungus is not identical with that produced by muscarine.

From another poisonous fungus, *Amanita phalloides*, Kobert<sup>2</sup> obtained a toxalbumin which he calls **phallin**. This substance is a blood-poison which disintegrates the red corpuscles, liberates fibrin-ferment, and causes the formation of thrombi. Fatty changes—especially in the liver—and the formation of multiple ecchymoses ensue, the train of symptoms closely resembling those due to acute phosphorus poisoning, the mucous membrane of the intestinal canal is injected, and the urine may contain hæmoglobin. Ford<sup>3</sup> denies that the hæmolyisin obtained by Kobert from *A. phalloides* is a proteid substance. He believes it to be a glucoside, and holds that the lesions produced by *A. phalloides* are to be attributed entirely to the amanitotoxin, which is the active principle of the fungus. In a case recorded by Handford<sup>4</sup> a man, aged thirty-two, ate about a quarter of a pound of cooked *A. phalloides*. In nine and a half hours he experienced a sense of weight and constriction in the chest, and pain in the bowels, he subsequently vomited and was purged, profuse sweating, dimness of vision, and headache occurred. When seen, about twenty-four hours after eating the fungi, the pulse was 92, small, scarcely to be felt at the wrist, and the respirations, of a sighing character, were 17 to the minute, the pupils were normal, the patient complained of pain in the abdomen, he was drowsy, passed very little urine, and became delirious, death took place on the third day. At the necropsy punctiform ecchymoses were found on the lungs and under the pericardium, the liver was in an advanced stage of fatty degeneration, the mucous membrane of the stomach was much congested and presented numerous points of capillary hæmorrhages and small superficial erosions, the whole of the intestines were slightly congested. A daughter of the deceased, who ate part of one of the fungi along with her father, vomited, and was purged, but had no abdominal pain, she died in twenty-nine hours. At the necropsy no signs of gastro-enteritis were found. Tappeiner<sup>5</sup> describes some cases of poisoning by *Amanita phalloides* in which neither the symptoms nor the post-mortem appearances indicated the occurrence of any solvent action on the blood corpuscles. The symptoms were either like those of cholera—as many as sixty to eighty stools in twenty-four hours—without important cerebral symptoms, or else they were of a nervous type with but slight intestinal disturbance—headache, somnolence, delirium, twitching of the muscles and

<sup>1</sup> Arch. f. exp. Path., 1903.

<sup>2</sup> Petersb. med. Wochenschr., 1891.

<sup>3</sup> Münchener med. Wochenschr., 1895.

<sup>4</sup> Brit. Med. Journ., 1906.

<sup>5</sup> The Lancet, 1886.

general convulsions, in some instances the pupils were dilated. Two of the cases in which the nerve symptoms were prominent died, they suffered from neither jaundice, hepatic pain, nor anuria. At the autopsy the intestinal mucous membrane was found to be slightly affected only, but there were small ecchymoses in various organs, and the kidneys and liver showed such advanced fatty changes that the latter resembled a phosphorus-liver. In one case the liver contained 68.9, and in another 53.6 per cent of fat. The heart was also fatty. Struble<sup>1</sup> relates eight cases of which three were fatal, of poisoning by *A. phalloides*. The symptoms began, in from nine to eighteen hours, by vomiting, without any indication of gastric irritation, followed by cardiac failure, collapse, and diarrhœa. There were no cerebral symptoms nor coma. In the three fatal cases death occurred in from forty-eight to fifty-four hours.

The *Helvella esculenta* is poisonous owing to the presence in it of helvellic acid, which acts much in the same way as phallin.

**Treatment of Poisoning by Fungi.**—Evacuate the stomach with an emetic and the bowels with castor oil, and then treat the symptoms. Atropine is recommended as an antidote. In poisoning by muscarine, it acts as nearly like a true physiological antidote as any antagonist well can do, unfortunately, in poisoning by fungi, even by the fly-fungus, its antagonism is less efficacious, still, it should be tried, especially if the symptoms partake of the muscarine type. Warmth and stimulants will probably be required, and morphine if the gastro-enteric symptoms prevail. In suspected cases, the motions should be carefully examined for spores.

**Post-mortem Appearances.**—The necropsies of the cases already quoted illustrate the salient post-mortem indications— inflammation of the gastro-intestinal mucous membrane, with hæmorrhagic spots and erosions, punctiform sub-pleural and sub-pericardial hæmorrhages, and not unfrequently signs of fatty changes in the solid viscera, especially in the liver, comprise the most important appearances. The fatty changes scarcely seem to have received sufficient attention. Maschka, Husemann, and Boudier<sup>2</sup> observed them many years ago both in animals and in the human subject, and since then numerous cases have occurred in which fatty liver has been recorded as one of the post-mortem signs of poisoning by fungi. It appears to be most common after poisoning by *Amanita muscaria* and *Amanita phalloides*, in Handford's and in Tappeiner's cases of poisoning by the last-named fungus, already quoted, the liver was in an advanced stage of fatty degeneration. Muller<sup>3</sup> examined the body of a woman who was found dead four days after having eaten part of a fly-fungus. The heart, kidneys, and liver all showed fatty changes, the liver especially presented such a typical "phosphorus-liver" appearance as to give rise to doubts whether death had been caused by phosphorus or by the fungus.

## BEANS.

Any kind of vegetable food which has undergone the initial changes of decomposition may cause toxic symptoms. A boy aged fourteen years ate eight or ten uncooked haricot beans, the following day he had severe headache and abdominal pain without vomiting or purging, the tongue was thickly coated and there was great thirst, but there was no temperature. The patient became mildly delirious and was very ill for three days, recovery took place slowly. The rest of the beans, cooked in the usual way, were eaten by the

<sup>1</sup> *American Med News*, 1899.

<sup>2</sup> *Des Champignons*, 1868.

<sup>3</sup> *Vierteljahrsschr. f. ger. Med.*, 1890.



family without ill effects. Fischer<sup>1</sup> records an outbreak of poisoning by beans which had been tinned, and were subsequently eaten in the form of salad. Twenty-four persons ate of the salad, and twenty-one of them were made ill, of these, eleven died. The incubation period was never less than twenty-four hours nor more than forty-eight. The symptoms comprised nausea, constipation, feebleness, double vision, ptosis, difficulty of speech, quick pulse (150 in one case), cyanosis, and death from respiratory paralysis in from two to eleven days. The symptoms resembled those of the neuroparalytic type of food poisoning, there was neither vomiting nor diarrhoea. Consciousness continued to the end. The post-mortem signs, apart from those due to asphyxia, comprised hyperæmia of and extravasations of blood into the mucous membrane of the lower intestinal tract, the stomach and the upper tract were unaffected. In some remains of the beans a bacillus identical with the *B. botulinus* of Van Ermengen was found, for the first time in a vegetable medium. The toxic agent was weakened or destroyed by the heat of cooking. Rolly<sup>2</sup> records an extensive outbreak of bean poisoning in which two hundred and fifty persons were affected with gastric disorder about four hours after eating some beans. *B. coli* and *B. paratyphi* were found in the beans, but not in the fæces of the sufferers. The outbreak was attributed to the toxine produced by the paratyphus bacillus. This toxine was not destroyed by moderate heat.

## CHAPTER XXXVI

## ANIMAL POISONS AND POISONING BY FOOD.

## CANTHARIDES.

**Cantharides**, or Spanish flies, contain an active principle or acid—*cantharidin* partly free and partly combined with organic and inorganic bases. It is insoluble in water, slightly soluble in cold alcohol, and more freely so in hot alcohol, fixed oils, ether, and chloroform, when combined with bases, its solubility in these solvents, respectively, is the converse of that which obtains when it is in the free state. Cantharidin is eliminated in the urine and fæces.

**Symptoms.** When taken internally in poisonous doses cantharides causes a burning pain in the throat, which quickly extends to the stomach, with difficulty in swallowing, intense thirst, salivation, swelling of the salivary glands, and vesication of those parts of the digestive tract with which it first comes in contact. Nausea and vomiting occur, the rejected matter containing shreds of membrane and probably blood, diarrhoea, with tenesmus, may follow. Pain in the lumbar region, accompanied by strangury and irritation of the urethra, is almost invariable, the urine contains albumin and occasionally blood. In severe cases collapse, coma, and convulsions may lead to death, which is usually due to paralysis of the respiratory centres. Womack<sup>3</sup> describes a case in which there was a reddish-bronze discoloration of the whole surface of the body, with a dark almost black patch extending across the nose on to both cheeks. The mucous membrane of the mouth was similarly suffused.

<sup>1</sup> *Zeitschr. f. klin. Med.*, 1906.<sup>2</sup> *Munchener med. Wochenschr.*, 1906.<sup>3</sup> *Brit. Med. Journ.*, 1911.

The hue became deeper before death. In another case, which recovered, there was a general yellowing of the skin. Both these cases were admitted to hospital for abortion.

When cantharides is criminally administered it is not with homicidal intent, the object is either to excite the sexual feeling or to procure abortion. In more than one instance death has accidentally resulted from a delirious patient eating a blister which was applied to his head. Cantharides is harmless to fowls, if, after being fed with it, a fowl is eaten by a human being, symptoms of cantharides poisoning are produced. Severe symptoms of poisoning have occurred from the external use of cantharides.

**Fatal Dose.**—The smallest recorded fatal dose is twenty-four grains of powdered cantharides, recovery has occurred after one drachm. One ounce of the tincture has caused death, recovery has followed after six ounces were swallowed. Very small doses may produce severe poisonous effects. Sedwick<sup>1</sup> records the case of a girl, thirteen and a half years old, to whom one Spanish fly was given in a tart, in half an hour giddiness, pain between the shoulders, and a burning sensation in the throat were experienced. The following morning, the abdomen was distended, there was strangury, and the vulva was swollen and irritable. The patient complained of a strong disagreeable odour in the nostrils, and vomited half a pint of blood, for three days there was occasional vomiting of blood, recovery took place. About seventy-five centigrammes (11½ grains) of cantharidin caused the death of a man of seventy in twelve or fourteen hours.<sup>2</sup>

**Treatment.**—Evacuate the stomach, and, if possible, wash it well out. Demulcents and morphine, with warm baths or fomentations, will be required. Fatty or oily substances must not be given.

**Post-mortem Appearances.**—Indications of inflammation are usually present in the mouth and along the digestive canal, they may diminish in intensity after the first part of the intestines is reached, or they may be continued as far as the rectum. Excoriation and ulceration of the mucous membrane, with swelling and softening, will probably be visible, producing in some cases the appearance of a raw blood-stained or purulent surface deprived of all trace of epithelium. If the powdered insect has been swallowed, bright, glistening particles are generally to be seen on the mucous or raw surface, especially of the intestines, in such cases, examination of the digestive tract with a lens should never be omitted. If death has occurred very shortly after the poison was swallowed, the changes in the stomach and bowels may be less pronounced. The kidneys are usually large, red, and gorged with blood, the epithelial cells of the glomeruli are swollen, softened, and detached, frequently blocking up the tubules, the internal surface of the bladder is injected, and is often ecchymosed, the mucous membrane of the urethra is also injected. The spleen has been found enlarged.

**Chemical Analysis.**—If the solid poison has been swallowed, the mucous membrane of the stomach and intestines should be scraped with the edge of a piece of glass and the scrapings distributed in water, by means of alternate agitation and decantation, fragments of the shining wing cases, which are easy of recognition, may be separated for microscopical and chemical examination. In order to obtain an extract of cantharidin from the tissues it will probably be necessary first to free the cantharidin from combination, either by simply acidulating with sulphuric acid or by Dragendorff's method, after which it may be dissolved out by shaking with chloroform. Dragendorff's method consists in boiling the organic mixture with potash and water, filtering, adding sulphuric acid to the filtrate to

<sup>1</sup> *Med Times and Gaz*, 1864.

<sup>2</sup> *Annales d'Hygiène*, 1892.

liberate the cantharidin from the potash, and then boiling the filtrate with four times its volume of alcohol, after cooling, the alcoholic solution is filtered, the alcohol evaporated, and the residue extracted with chloroform, on evaporation of the chloroform, a portion of the final residue may be taken up with a little oil

**Tests.**—A morsel of cotton-wool saturated with the only mixture obtained from the chloroform extract, and retained for some hours in contact with the skin on the arm or the breast, will raise a blister if even but a very minute quantity of cantharidin is present. An aqueous solution of cantharidin in combination with potash or soda yields a green precipitate with copper sulphate, and a red precipitate with cobalt sulphate.

Cantharides resist putrefaction for a long time

### POISONING BY FOOD.

It was formerly believed that the appearance of toxic symptoms following the ingestion of food was due to the development of poisonous principles in the food caused by putrefactive changes. The symptoms were attributed to the action of animal alkaloids or ptomaines, and were spoken of as "ptomaine poisoning." At a later date it became recognised that many of the outbreaks of food poisoning were really due to bacterial infection, and it is now believed that almost all such outbreaks are caused in this way, and that ptomaines play only a small part, if any, in food poisoning. Savage<sup>1</sup> has recently reviewed the evidence held to support the toxicity of ptomaines. He points out that the belief in their poisonous properties was almost entirely founded upon the results of *inoculation* experiments in animals. It is now known, however, that many substances, such as snake venoms and products of pathogenic organisms, are intensely toxic when introduced under the skin, but when taken by the mouth produce symptoms only when enormous doses are swallowed. Savage has failed to come across any direct evidence that feeding with ptomaines prepared from putrefying meat has produced symptoms of food poisoning. Nor is the evidence any stronger that the symptoms are due to products of putrefaction other than ptomaines or to toxins produced by the activities of putrefactive bacilli. Experiments made by feeding animals on highly putrid meat failed to produce symptoms of poisoning. On the other hand, subcutaneous injection of but one cubic centimetre of washings from putrid meat proved fatal.

Food may become poisonous in the following ways. —(1) Pathogenic organisms may be present in the milk or in the flesh of an animal when slaughtered, or may be introduced into the food during the process of preparing it for the table. The conveyance of tuberculosis, diphtheria, typhoid, etc., by these means belongs to the domain of public health. Most of the acute outbreaks of food poisoning have been due to infection with a bacillus of the Gærtner group. Infection has often been conveyed by eating the meat of calves suffering from septic infection of the navel or cows which have been infected after calving or display infection of the udder. In other cases diseased sheep or pigs have been responsible. The risk of transmission of infection by diseased meat is lessened by cooking, but this is not an infallible protection. Many outbreaks have been due to food eaten in an uncooked or incompletely cooked state, such as smoked ham, pies, and milk. Cooks and other persons who handle food may infect it with pathogenic organisms if they are themselves suffering from an infection. An interesting case of food poisoning in which infection was conveyed by the

<sup>1</sup> *Food Poisoning and Food Infections*, 1921.

person who handled the food was investigated recently in South London<sup>1</sup>. On Saturday a stew of steak and liver was prepared and consumed by a family, the gravy being saved until the following day and warmed up with a Yorkshire pudding. On the previous Thursday the landlady, who prepared the food, was taken ill, and the climax of her illness was reached on Saturday night, though she continued to attend to her household duties. The vehicle of infection was the gravy from the liver. All the nine persons who ate this displayed gastro-intestinal symptoms and two died. A bacillus of the Gærtner group was present in the organs of the deceased and in the blood of those who survived.

(2) Sound food may become contaminated with chemical poisons, either derived from the container or introduced as adulterants or preservatives. Instances of poisoning by arsenic and lead have been given in the preceding pages. There is no evidence that tin, which is so largely used for holding food, has any toxic effect. The effects of preservatives and adulterants in foodstuffs are dealt with in books on public health.

(3) Food may be poisonous to certain persons owing to a special sensitiveness or idiosyncrasy. It has long been known that certain persons may exhibit toxic symptoms after eating some form of food quite innocuous to normal individuals. Cases are recorded of persons in whom a small quantity of white of egg invariably produced urticaria, vomiting, increased respiration, and even coma. Other foods towards which such abnormal sensitiveness has been shown are blackberries, almonds, tomatoes, and cheese. These symptoms are now regarded as instances of anaphylactic shock.

**Poisoning by Meat.**—As already stated, this is most often due to infection by the bacillus of Gærtner. A small group are due to infection with the bacillus botulinus. It is important to notice that the food may display no indications of any sort either in taste, smell, or appearance, that it is different from sound food. The incubation period varies from a half to forty hours or more. The onset is usually abrupt, and the symptoms are those of marked gastro-intestinal irritation with involvement of the nervous system. Vomiting with severe diarrhoea, colic, and pain in the abdomen, are present. Skin rashes, such as erythema, urticaria, herpes, and purpura, may occur. Nervous symptoms include numbness and cramps in the limbs, photophobia, and delirium. Signs of collapse with cold sweats and even rigors may precede death. The case mortality rate, however, is not high. In 112 British outbreaks tabulated by Savage, comprising 6 190 cases, there were 94 deaths, a case mortality of 1.5 per cent. Outbreaks are more frequent during the summer than the winter months. The post-mortem signs are those of gastro-enteritis.

**Botulism** is a form of food poisoning rare in this country. The toxins of the organism responsible, *B. botulinus*, are destroyed by efficient cooking. Hence the condition has been found most commonly after taking food which has not been cooked, or has been insufficiently cooked, such as sausages, smoked ham, salad, or beans which have been merely rinsed and warmed. The symptoms, which appear in from 12 to 36 hours, affect chiefly the central nervous system. They comprise thirst, a feeling of constriction of the throat, obstinate constipation, amaurosis, paralysis of accommodation, ptosis, diplopia, etc., muscular weakness, and in fatal cases paralysis of the respiratory and cardiac systems. Vomiting and diarrhoea may occur, but are frequently absent. The case mortality may be as high as 30 to 50 per cent.

In the diagnosis, differentiation has to be made from polio-encephalitis, bulbar paralysis, and the various ophthalmoplegias.

<sup>1</sup> *The Lancet*, August, 1920.

## POISONING BY FISH.

Certain kinds of fish of the species *tetrodon* (*fuga*) found in Japanese waters, and several others mostly of tropical origin are inherently poisonous, mackerel, carp, barbel, and herrings are liable to become poisonous at times, some of them being specially prone to develop poisonous properties after they are dead, mackerel, for example, speedily becomes unfit for food, herrings also, especially when they are not "gutted" immediately after being caught. Poisoning has often been caused by caviare and by the roe of herrings and of other fish. Dried salt cod-fish and preserved anchovies have proved poisonous, a stale red herring caused the death of a man from gastro-enteritis. Shell fish excepted, mackerel is probably answerable for the greatest number of cases of fish poisoning in this country. The symptoms are usually gastro-enteric, Addinsell<sup>1</sup> saw a man who, after eating fresh mackerel, developed gastro-enteritis followed by fever with the formation of a rectal abscess, the patient recovered after a long illness. As with meat, the toxin may be limited to a part or parts of the fish.—A man ate some mackerel which showed signs of decomposition about the gills, he was taken ill with acute gastritis and delirium, and died, his wife also ate of the same fish without ill effects. It turned out that the man ate of the parts nearest the head where the putrefactive processes were most evident, the tail end fell to the lot of his wife. Poisonous fish, however, do not always attack the intestinal tract. Edwards<sup>2</sup> relates the case of a woman who, after eating some ray fish, suffered from a swelling of the face and tongue, the latter filling the whole mouth, a burning sensation was felt in the hands and back of the head, with cold feet, thirst, dyspnœa, slight convulsions, and intense itching of the skin, she had no pain, and she soon recovered. The symptoms were apparently due to some toxin which acted chiefly on the nervous system.

**Tinned fish** on many occasions has caused poisoning. Six persons ate tinned salmon at supper, early the following morning they were seized with violent pain in the stomach, vomiting, profuse diarrhœa, pain in the head, thirst, a temperature of 102° to 103° F., and a pulse-rate from 110 to 160 in the minute, one patient became semi-unconscious, with a temperature of 104° F., the pulse was almost imperceptible, the skin was cold and clammy, and pupils were widely dilated, death ensued. At the necropsy the brain was found superficially congested, and parts of the stomach and the intestines were so deeply inflamed as to be almost gangrenous.<sup>3</sup> Stevenson<sup>4</sup> records the case of a man, aged twenty-one, who ate six sardines for breakfast, a few hours after he complained of feeling unwell, and vomited. Next morning there was slight pain in the stomach, the abdomen was tense but not enlarged and the patient was perspiring, shortly after noon collapse rapidly set in, and death occurred almost immediately. At the necropsy made the following day the features were found bloated so as to be unrecognisable, although the weather was cold (47° F.), blood-stained fluid exuded from mouth, nostrils, and ears except the hands and feet, the whole body was emphysematous, and there were large bullæ on the buttocks. The abdomen and the bladder were distended with gas, the mucous membrane of the stomach and intestines was emphysematous, the liver was cavernous and friable, and along with the kidneys and bladder was hyperæmic, the large intestine was normal, and contained solid fæcal

<sup>1</sup> *The Lancet*, 1889<sup>2</sup> *Brit Med Journ*, 1884<sup>3</sup> *Brit Med Journ*, 1891<sup>4</sup> *Ibid*, 1892

matter These extremely rapid putrefactive changes were probably due to the presence of micro-organisms in the fish

**Shell Fish.**—Some kinds of shell fish may become contaminated with pathogenic micro-organisms, and may cause a true infection in human beings who eat them, this has been observed in oysters and mussels (eaten raw) which have grown or have been placed in water polluted with sewage containing typhoid germs—e.g., in the estuaries of rivers near towns Experiments by Wood<sup>1</sup> tend to show that the bacilli of typhoid fever can probably live in sea-water for at least two months, Boyce and Herdman<sup>2</sup> found them on the twenty-first day On the other hand, Frankland<sup>3</sup> and Cassedebat<sup>4</sup> state that they are rapidly destroyed by sea-water Conn<sup>5</sup> investigated the etiology of an epidemic of typhoid fever that occurred in a college, twenty-six persons, of whom four died, being attacked He traced the infection to some oysters which had been grown in deep water and had been subsequently deposited at the mouth of a fresh-water creek to “fatten” them—oysters thus treated absorb an amount of fresh water which gives them a plump appearance Three hundred feet from the oyster bed the mouth of a drain was found which came from a house where there were two cases of typhoid fever Numerous instances in which oysters have communicated typhoid fever are recorded by Broadbent,<sup>6</sup> News-holme,<sup>7</sup> and others

**Mussel poisoning** is due to the formation of a toxin whilst the fish (*Mytilus edulis*) is still alive It was formerly believed that mussels owed their poisonous properties to the presence in them of copper derived from ships' bottoms, or from copper-covered fixtures in harbours, to a special disease which the fish laboured under, to the presence of specimens of a poisonous species along with the edible fish, to incipient putrefaction taking place after the harmless fish were removed from the water, and to various other conditions which were assumed in default of positive knowledge Brieger<sup>8</sup> was the first to succeed in isolating from poisonous mussels a basic product in a condition sufficiently pure to admit of ultimate analysis A number of cases of mussel poisoning, several of which were fatal occurred at Wilhelmshaven in 1885, from a quantity of the noxious mussels Brieger obtained a poisonous base which he called *mytilotoxin*, to which he ascribed the formula  $C_8H_{15}NO_2$  other bases, among which is *betaine*, were also present

The conditions which lead to the development of this toxin in mussels are—stagnant water, or water which is not in free communication with the sea, or water which is contaminated with sewage, or other organic matter undergoing decomposition It is not essential that the water should contain noxious matter, mere absence of freshness is sufficient to interfere with the metabolism of the mussels to such an extent as to lead to abnormal changes in their tissues, which result in the formation of a toxin during life if mussels which have thus become poisonous are placed in water in free communication with the sea, they rapidly lose their poisonous properties Poisonous mussels have invariably been obtained from harbours, docks, the mouths of rivers, and other places where either a deficiency of tidal interchange takes place, or where the water is contaminated with decomposing organic matter

**Symptoms.**—The mild and common form of mussel poisoning is characterised

<sup>1</sup> *Brit Med Journ*, 1896

<sup>2</sup> *Rep of the Brit Assoc*, 1896

<sup>3</sup> *Proc of the Royal Soc*, 1894

<sup>4</sup> *Revue d'Hygiene*, 1894

<sup>5</sup> *Medical Record*, 1894

<sup>6</sup> *Brit Med Journ*, 1895

<sup>7</sup> *Ibid*, 1896

<sup>8</sup> *Ueber Ptomaine*, *Dritter Theil*, 1886

by the appearance of an exanthematous, or urticarial eruption on the body, which may be associated with a feeling of oppression in the chest and difficulty of breathing, in the severer forms gastro-intestinal disturbance occurs, and, in the most dangerous of all, paralysis

The following cases illustrate the etiology and the symptoms of fatal mussel poisoning. Permevan<sup>1</sup> relates the case of a man aged forty who ate uncooked a large quantity of mussels obtained from the bottom of a graving dock. When seen a few hours after, he was absolutely unconscious, the face was livid, the pulse almost imperceptible, the pupils were widely dilated and inactive, and the limbs were flaccid from complete paralysis, he took gasping breaths about once or twice in the minute, there was no vomiting nor purging, and the temperature did not materially fall until the circulation failed. Neither stimulants, atropine, strychnine, nor artificial respiration evoked any attempts at natural respiration, and death took place in about twelve hours after the mussels were eaten, the heart continued to beat for many hours after voluntary respiration had ceased. In another case, reported by Cameron,<sup>2</sup> a woman and five children ate some mussels which had been gathered from a sheet of water to which the sea had access, and into which fresh water and some sewage flowed. In twenty minutes symptoms of poisoning commenced in the form of prickly pains in the hands, in less than one hour one child died, in two hours the mother and three other of the children were dead. They suffered from violent vomiting, hyspnœa, swelling and lividity of the face and spasms, they appeared to have died asphyxiated. One child and a maid who ate but few of the fish recovered. Cameron found an alkaloidal substance, and McWeeney<sup>3</sup> found bacteria in some mussels from the same source. A very rapid death from mussel poisoning is recorded by Hill.<sup>4</sup> A man aged forty-nine ate some mussels, very soon after his face became fiery red and he felt an itching sensation in his hands and legs, he rapidly became worse and died in about an hour after the symptoms commenced. At the autopsy the gastric mucous membrane showed no signs of inflammation, nor was there any obvious cause of death.

Other shell fish besides mussels may develop toxins. Cameron relates a case in which ten out of twelve persons, who lunched together, ate some oysters, nine out of the ten were attacked with nausea, vomiting, diarrhœa, abdominal pain, and prostration, they all recovered. The oysters had been grown in a place to which sewage had access. Casey<sup>5</sup> records a fatal case, a man aged thirty-two ate eight oysters, remarking at the time that one was bad. About twelve hours after, he began with pain in the back and in the abdomen and vomited frequently, but there was no diarrhœa, he became collapsed and died forty-one hours after eating the oysters. The stomach was found to be darkly congested and the peritoneum was thickly studded with flakes of lymph. In other instances oysters have caused symptoms of a purely neurotic type. Brosch<sup>6</sup> relates such a case, fatal in twenty-two hours, in which vomiting, giddiness, paralysis of the muscles of the pharynx and bladder, with ptosis and loss of power of accommodation occurred, death being due to paralysis of the muscles of respiration. After death the brain was found œdematous and there were punctiform hæmorrhages in the cerebellum, and the lower dorsal and upper lumbar regions of the cord and also on the pericardium and pleuræ, the spleen was enlarged and there were fatty changes in the liver. No

<sup>1</sup> *The Lancet*, 1888

<sup>2</sup> *Brit Med Journ*, 1890

<sup>3</sup> *Ibid*, 1890

<sup>4</sup> *Ibid*, 1895

<sup>5</sup> *Brit Med. Journ*, 1894

<sup>6</sup> *Wiener klin Wochenschr*, 1896

micro-organisms were found, but ptomaine-like bodies were detected in the viscera

**Treatment.** Evacuate the stomach with an emetic, and the bowels with an aperient, unless both vomiting and purging have already emptied the digestive canal, then give stimulants, apply external warmth and friction, and perform artificial respiration if required. Atropine has been recommended. Morphine may be advisable if the purging and abdominal pain are excessive.

**Post-mortem appearances** are not characteristic, there may be indications of gastro-intestinal irritation.

**Crabs.** Poisoning by the larger shell-fish may give rise to pronounced post-mortem appearances. A young man, previously healthy, was admitted into hospital suffering from violent abdominal pain, vomiting, diarrhoea, and cramps, which came on a few hours after eating two small crabs that were "not quite fresh." The acute symptoms gradually subsided, but occasional vomiting persisted. He could not take food, and he progressively emaciated and became weaker, and eventually died from inanition in seven weeks. On section, the stomach was found to be reduced to a small cavity in the midst of adhesions, it was devoid of mucous membrane and could only be identified by its anatomical surroundings. The intestines were not materially affected.

### POISONING BY MILK AND CHEESE.

**Milk** may become contaminated with pathogenic micro-organisms, and may thus cause infection in human beings who drink it. Gaffky<sup>1</sup> describes how some milk from a cow which had severe enteritis made three persons ill with stupor, delirium, high fever, vomiting, albuminuria, and copious motions, some of which contained blood. Inherently the milk was free from bacteria, the infection having been caused by contamination of a portion of it with some of the cow's dejections, bacilli of the same kind were found both in the motions of the cow and also in those of the patients. Niven<sup>2</sup> reports an outbreak of milk-infection which occurred in Manchester in 1894, the milk which caused the mischief was supplied by one dealer, and in it Delépine found a streptococcus and a bacillus of the type of *B. coli commune*. Welply<sup>3</sup> relates several instances in which typhoid fever was communicated by milk, either on account of the cans having been washed with contaminated water, or in some instances by direct contamination of the milk from the hands of dairymaids who, at the time, were nursing some patients suffering from enterica. Tower<sup>4</sup> enumerates a number of diseases in cows which may make their milk dangerous to human beings. From experiments made by Flugge<sup>5</sup> it appears that boiling is not sufficient invariably to sterilise milk, spores are not destroyed by being boiled for three-quarters of an hour or more, and some anaerobic bacteria resist still longer. Sterling<sup>6</sup> supports Flugge's views as to the difficulty of sterilising milk, especially as regards certain peptonising bacteria.

Kerr and Hutchens<sup>7</sup> described an extensive outbreak of poisoning caused by drinking infected milk, which produced serious symptoms in at least 523 persons living in and around Newcastle. The period of incubation varied from four to thirty-nine hours, the majority being less than eighteen hours.

<sup>1</sup> *Deutsche med. Wochenschr.*, 1892

<sup>4</sup> *Medical News*, 1891

<sup>2</sup> *The Lancet*, 1895

<sup>5</sup> *Zeitschr. f. Hygiene u. Infektionskrankheiten*, 1894

<sup>3</sup> *Ibid.*, 1894

<sup>6</sup> *Medycyna*, 1895

<sup>7</sup> *Proc. Roy. Soc. of Med.*, 1914



The symptoms displayed were severe vomiting, diarrhœa, and prostration, accompanied by abdominal pain and high temperature. No fatal cases occurred. The infection was traced to a cow which had recently calved. She displayed illness for a couple of days and her milk was diminished in quantity, and was noticed by the milkman to be "rather thick like strippings." Nevertheless it was mixed with the rest for sale. On the following morning, the cow was found dead in her stall. Gärtner's bacillus was isolated from the spleen, mesenteric glands, uterus, intestine and milk of the cow, and from the stools of seven of the persons who had been affected by the outbreak.

Milk has caused severe poisoning owing to the occurrence in it of chemical changes. Newton and Wallace<sup>1</sup> gave an instance in which a number of people living in two hotels were suddenly taken ill four hours after supper with symptoms of gastro-intestinal irritation—nausea, vomiting, cramps, and collapse, a few had diarrhœa. The following week a second series of cases, of precisely the same nature, occurred in another hotel, and the symptoms were traced to the milk, but no added poison was found in it. Further investigation showed that the cows from whence the milk was derived were healthy, but that after milking, the milk was at once placed in cans and carted eight miles during the warmest part of the day, and in a hot season of the year, the method usually adopted being to allow the milk to cool in shallow open vessels surrounded with cold water or ice previous to transport. Chemical examination of the suspected milk revealed the presence of a substance—identical with Vaughan's tyrotoxin (*vide infra*)—which produced symptoms of poisoning in a cat. Leeds<sup>2</sup> states that **condensed milk** may undergo changes due to bacterial ferments present in the milk before condensation, these changes are shown by a tendency either to solidification of the milk, or to putrefactive processes. Ice creams have often proved toxic from contamination with pathogenic micro-organisms owing to the insanitary conditions under which the creams are frequently made. In some instances children who have eaten poisonous creams are attacked with symptoms resembling those of meningitis—restlessness, apathy, insensibility and retraction of the head—which usually subside in forty-eight hours.

**Cheese.**—Numerous instances of poisoning by cheese have occurred, especially in Germany and America. The symptoms which appeared in a number of cases (nearly three hundred) that happened in America in 1884 and 1885 were as follows—Violent vomiting and diarrhœa, pain in the stomach and cramp in the legs, the tongue at first coated white, was later red and dry, the pulse was weak and irregular, and the face pale and cyanotic. None of the cases were fatal. The cheese which caused the symptoms was not old nor decayed, Vaughan<sup>3</sup> found that it possessed the characteristic of instantly and intensely reddening litmus-paper, good cheese when new slightly reddens litmus-paper, but does not produce immediate and pronounced change of colour.

By extracting the poisonous cheese with water, adding alkali, and shaking out with ether, Vaughan obtained needle-shaped crystals, which had a distinct toxic action, and to which he gave the name *tyrotoxin*. The substance is not an alkaloid nor does it respond to the alkaloidal group-tests, it is soluble in water, alcohol, ether, and chloroform. It appears to be due to the action of micro-organisms present in the milk from which the cheese is prepared. The chemical reactions and, to some extent, the physiological action of tyrotoxin resemble those of diazo benzene. Vaughan thinks that the two substances are

<sup>1</sup> *Medical News*, 1886

<sup>2</sup> *Amer Journ Med Sc*, 1895

<sup>3</sup> *The Practitioner*, 1887

identical. Subsequently Vaughan<sup>1</sup> found a toxalbumose in some cheese in which no tyrotoxicon was present. Old decayed cheese yields an alkaline reaction, and has frequently given rise to colic, diarrhœa, dizziness, diplopia, precordial pain, and collapse. Brieger obtained trimethylamine from decayed cheese.

Probably most cases of poisoning by cheese are due to the presence of toxins generated by bacterial activity, and it is likely that investigation by modern methods will show that a bacillus of the Gærtner group is most often responsible.

<sup>1</sup> *Phil Med and Surg Repr*, 1895

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